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A thesis submitted by Sean Doherty B.Sc. in accordance with the requirements for the degree of Doctor of Philosophy in the University of Durham.

Department of Biological Sciences, August 2000

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2 0 MAR 2001

For my Dad

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#### ABSTRACT

#### Apoplastic proteins, enzymes and radicals.

The soluble and readily extractable part of the plant extracellular matrix has been termed, the apoplast and contains a wide range of components such as, complex carbohydrates, structural proteins, enzymes and radicals that are known to be responsive to stress and developmental pressures.

This thesis describes the development of a technique for the selective enrichment of apoplastic components for a range of subsequent analyses. Using this technique a number of apoplastic proteins were N-terminally sequenced and revealed 2 cell wall related enzymes, an antifungal protein and 3 auxin-binding/germin-like proteins. This technique also provided a novel approach to the further study of auxin-binding proteins via the use of affinity chromatography at their putative site of action, the apoplast. Three potential auxin-binding proteins were identified.

Many attempts were made to subject the material extracted from the apoplast to the highly resolving technique of 2-dimensional electrophoresis, and during the process two unusual 2D systems were developed. These systems could be run in a small format that permitted very rapid analysis and/or using an in-gel loading strategy to subject up to 500µg of protein to 2D separation therefore permitting N-terminal sequencing from single 2D gels. Unfortunately 2D separation of apoplastic proteins was never fully achieved within the time frame of this study due to the vast degree of heterogenity present in the sample material. It did however demonstrate the very complex nature of apoplastic components.

A series of experiments revealed that the tobacco leaf apoplast contained compartment specific antioxidant enzymes, some of which share physical characteristics with similar enzymes from other species. The activity of these enzymes altered in response to stress and according to the developmental age of the tissue. The reduced activity of these enzymes directly correlated to the degree of oxidative modification of apoplastic proteins illustrating that these enzymes are important in the detoxification of apoplastic radicals.

Follow on experiments following the apoplastic generation of the superoxide anion and nitric oxide from impact stressed potato tuber tissue showed that radicals play important roles in the responses of plant tissue to stress, and show the first involvement of nitric oxide in plants in response to abiotic stress.

#### STATEMENT

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Ser Dhy

Sean Doherty

#### MEMORANDUM

# Parts of this work have been submitted for publication in the following peer reviewed international journals:

#### Doherty, S. & Croy, R.R.D.

Tobacco leaf apoplastic superoxide dismutases and peroxidases. *Plant, Cell and Environment Submitted September 2000; the manuscript is presented in Appendix 1.* 

#### Johnson, S.M., Doherty, S. & Croy, R.R.D.

Superoxide radical production in potato tubers: a novel biphasic response to mechanical stress. *Planta Submitted August 2000; the manuscript is presented in Appendix 1.* 

#### Doherty, S. & Croy, R.R.D.

A simple and rapid microtitre plate spectrophotometric assay for superoxide dismutase. *Pre-submission manuscript is presented in Appendix 1.* 

#### Parts of this work have been presented at the following International Conferences:

#### Doherty, S. Johnson, S.M., Shehab, G.M.G. & Croy, R.R.D. (1999)

Radical Responses to stress in potatoes. Presented at The 13<sup>th</sup> John Innes Symposium; 'Attack & Defence in Plant Disease'; Norwich, UK, 20-23<sup>rd</sup> July 1999.

#### Bowler, K., Bates, T. & Doherty, S. (1998)

Thermal tolerance in adult blowflies protects mitochondria from heat damage and involves production of heat shock proteins. Presented at the XIX Congress of the European Society from Comparative Physiology and Biochemistry; 'Cellular and Molecular Responses to Environmental Changes'; Turku, Finland, 23-26<sup>th</sup> August 1998

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#### ABBREVIATIONS

ABP	auxin binding protein
AGP	arabinogalactan protein
AOS	active oxygen species
Bis-acrylamide	bis (N,N'-methylene-bis-acrylamide)
BSA	bovine serum albumin
°C	degrees centrigrade
cm	centimetre
CMF	cellulose microfibril
2D	2-dimensional electrophoresis
dH <sub>2</sub> O	distilled water
DNPH	dinitrophenylhydrazine
DTT	dithiothreitol
EDTA	ethylenediamine tetra-acetic acid
q	gram
ĞalA	galacturonosyl acid
GRP	glycine-rich protein
GSH	reduced glutathione
GSSG	oxidised glutathione
GSNO	nitrosoglutathione
	hydrogen peroxide
	hydroxyphenylacetic acid
	hydroxproline-rich alvcoprotein
	intercellular fluid
	isoelectric focussing
	kilodalton
	litro
1 5.4	molar
	molocular weight
	molecular weight
MDH	
NaCl	sodium chioride
NADH	nictotinamide adenine dinucleotide
NADPH	nictotinamide adenine dinucleotide
NEDD	N-(1-napthyl)-ethylenediamine dinydrochloride
NO	nitric oxide
OH	hydroxyl radical
OONO	peroxynitrite radical
OONONOH	peroxynitrous acid
OONOCO <sub>2</sub>	nitrosoperoxocarbonate molecule
PAG-IEF	polyacrylamide gel isoelectric focusing phosphate
pl	isoelectric point
PGA	polygalacturonic acid
PMS	phenazine methosulphate
PR	pathogenesis-related
PRP	proline-rich protein
PVDF	polyvinylidene difluoride
QH <sub>2</sub> O	deionised water
RG	rhamnogalacturonan
SDS-PAGE	sodium dodecyl sulphate polyacrylamide gel electrophoresis
SULF	sulphanilamide
TE	total extract
Tris-HCl	tris(hydroxymethyl) methylamine hydrochloride
RUBISCO	ribulose bis-phosphate carboxylase
VI	vacuum infiltration
	volume for volume
	volume for weight
V/W XC	volation
	2 3-bis-12-methoxy-4-nitro-5-sulphonhenvll-2H-tetrazolium-5-
	carboxanilide

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#### **SECTION 1**

#### INTRODUCTION

#### 1.1 The Plant Extracellular Matrix

With growing knowledge the antiquated view of the plant extracellular matrix as a relatively inert, static and homogenous compartment has been radically revised to produce a concept of the extracellular matrix as a near extension of the cytosol, with a unique biochemistry and a complex dynamic structure. As an integrated structure, components in the extracellular matrix partly determine the development of the plant form, being essential for morphogenesis, organogenesis and therefore plant growth in its entirety. As would be expected of such a fundamental compartment of a plant, its structure, composition and biochemistry is dynamic during all stages of cell growth, maturation and differentiation.

Intracellular mechanisms exist to respond to temperature fluctuations, drought, changes in salinity, heavy metal presence and elevated levels of atmospheric pollutants. In tandem with intracellular changes, the extracellular matrix is also modified to enable the plant to cope with any such external environmental stress. In an analogous manner, adaptation is evident when a plant is wounded or undergoes pathogen attack. All adaptations are presumed to involve enhancing, restructuring or altering the composition of the extracellular matrix, to protect the plant and increase its overall chances of survival.

For the course of this introduction the plant extracellular matrix will be sub-divided into its two main discrete components, the cell wall and the apoplast. The cell wall is described in section 1.2 and the term apoplast will be fully defined in section 1.3. Figure 1.1 presents a highly simplified diagram to illustrate this sub-division.

#### 1.2 The Plant Cell Wall

The cell wall is well recognised as the principal structural element of the plant that imparts mechanical integrity, for controlling the expansion of cells during development and acting as the primary 'barrier' between the external environment and the intracellular milieu [Braam 1999].

Cellulose is the principal element of the cell wall; it is synthesised at the plasmamembrane and forms long crystals of several dozen glucan chains. Cellulose in





Intracellular Environment

#### Figure 1.1

# Schematic of the intracellular and external environments and the extracellular matrix.

Changes in the external environment can stress the plant and ultimately change both the intracellular environment and the extracellular matrix to cope with the imposed external stress. The plant extracellular matrix can be sub-divided into the plant cell wall, which comprises of 3 independent domains (see section 1.2) and the apoplast (see section 1.3).

conjunction with the heterogeneous xyloglucan group of polysaccharides provides the microfibrilar foundation of the cell wall that encases the cell and houses a diverse range of other non-cellulosic polysaccharides, proteins and enzymes. Indeed, several recent reports have altered the historical view of the cell wall as static and purely structural in function to a virtual continuum of the cytoskeleton [Lin *et al.* 1990; Wyatt & Carpita 1993; Zhu *et al.* 1993; Freshour *et al.* 1996].

Although the primary cell wall's of all plant species contain the fundamental cellulosic microfibrilar structure; other components are extremely diverse [Gibeaut & Carpita 1994]. Leaders in the field of plant cell wall biology, Carpita & Gibeaut [1993] have sub-divided the components that comprise the primary cell walls of plants into three independent, yet, interacting domains (see figure 1.1). Domain 1 comprises the basic structural elements of the plant cell wall, the cellulose-xyloglucan framework, which is embedded within a range of related gelatinous compounds that form the second domain, the pectin matrix. Both of these domains are comprised wholly of carbohydrates. The third independent domain consists of the proteinaecous elements of the plant cell wall.

Carpita & Gibeaut [1993] have also proposed that all plant cell walls can be divided into one of two broad categories; type I, which is typical of most flowering plants, and type II, which is seen primarily in the genus, Poaceae and some monocots. Throughout this research plants with type I cell walls have been utilised for study, consequently, this introduction will be limited to detailing the current consensus of a generalised type I cell wall. Furthermore, instead of constantly referring to the type I cell wall; it will simply be referred to as, the cell wall.

#### 1.2.1 Domains 1 & 2 : The plant cell wall polysaccharides

Cellulose and callose are the only known plant polysaccharides to be synthesised at the plasma-membrane [Delmer 1991]. All other polysaccharides are synthesised by the Golgi apparatus [Driouich *et al.* 1993]. Not a single synthase of the vast array of polysaccharides present in plant cell walls has ever been unequivocally identified because, all plant polysaccharide synthases appear to be membrane-spanning proteins which are uniquely sensitive to solubilisation with respect to their enzymatic activities [see Driouich *et al.* 1993 for a review]. Consequently, it is only very recently, with the advent of new methodologies for polysaccharide sequencing that the myriad of polysaccharides present in the cell wall, are beginning to catalogued and categorised [McCann *et al.* 1994; Watt *et al.* 1999]. Due to the highly heterogeneous nature of cell wall polysaccharide material and relatively scarce information regarding functionality, the description below will be limited to the cellulosic microfibrilar foundation (domain 1) and an overview of some of the more prevalent and well documented polysaccharides found

housed within this framework (domain 2). Figure 1.2 provides a reference for structural information regarding the major polysaccharides discussed within the text.

#### Domain 1 : The cellulose-xyloglucan framework

The cellulose-xyloglucan framework is a fibrilar network encasing the cell that accounts for approximately 50% of the primary cell wall mass. It forms the structural foundation of the cell wall and is composed of approximately 50% cellulose and 50% xyloglucan [Carpita & Gibeaut 1993].

Cellulose,  $(1\rightarrow 4)\beta$ -D-glucan chains of varying length, is synthesised at the plasmamembrane, by an as yet unidentified cellulose synthase complex and liberated into the extracellular matrix [Brett 2000]. Individual units of cellulose hydrogen bond to each other along their length to form paracrystalline assemblies of relatively huge cellulose microfibrils (CMFs) approximately 5-15nm wide [McCann *et al.* 1990]. Detailed electron microscopy studies have revealed that CMFs are wrapped around the cell like a skeletal cage, with individual fibres regularly spaced approximately 20 to 40 nm apart [McCann *et al.* 1990]. The exact arrangement of the CMF 'cage' alters according to cell and tissue type and developmental stage [Carpita & Gibeaut 1993] (see section 1.2.3).

Xyloglucans (XGs) are synthesised in the Golgi apparatus and secreted at the plasmamembrane surface by the secretory system. XGs have essentially the same glucan 'backbone' structure as cellulose, which comprise of linear chains of condensed  $(1 \rightarrow 4)\beta$ -D-glucan monomers approximately 200 nm in length [McCann et al. 1990]. The most consistent structural difference is that XGs have numerous  $\alpha$ -D-xylosyl units added at the O-6 position of the glucan residues within the backbone [Bacic et al. 1988]. However, XGs can be sub-categorised because of a bewildering array of potential tri-, hepta-, undeca- and many other oligomeric additions of monosaccharide units such as, β-Dgalactose,  $\alpha$ -L-arabinose and  $\alpha$ -L-fucose to the xylosyl branch of the glucan backbone [Bacic et al. 1988]. The effect of these multitudinous structural differences can only be guessed at, but what is certain is that XGs have a very distinct and different functionality to cellulose, especially with respect to hydrogen bonding characteristics. Computer simulations have predicted that XGs can assume a range of low energy conformations, all with the potential to hydrogen bond to exposed  $\beta$ -D-glucan chains in CMFs [Levy 1991]. Consequently, it has been postulated that XGs can hydrogen bond to one another and to CMFs thereby aiding in spanning, linking and locking the CMF 'cage' around the cell [McCann et al. 1992; Carpita & Gibeaut 1993]. Figure 1.3 presents a diagrammatic representation of the current consensus.

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#### Figure 1.2

#### Plant cell wall polysaccharides.

Structures of the key plant cell wall polysaccharides described within the text. Red arrows and text indicate sites and description of potential additions and changes to the rudimentary structure that generate a high degree of heterogeneity. Structures are derived from information provided in Carpita & Gibeaut [1993].



#### Figure 1.3

#### The 3 domains of the mature plant cell wall.

The diagram illustrates a single stratum of cellulose microfibrils (CMF's) in parallel arrangement, as they would be around a cell; the various CMF-linking molecules involved in structural integrity are presented in the legends. Several strata would form a full section of cell wall. This image has been adapted from Carpita & Gibeaut [1993].

There are many other non-cellulosic polysaccharides found in association with the cell wall, such as, galactoglucomannans, galactomannans,  $(1\rightarrow 3)\beta$ -D-glucans and glucutonoarabinoxylans, which also have the potential to bind to CMFs, but are found in much smaller amounts relative to XGs [Bacic *et al.* 1988; Maltby *et al.* 1979]. These polysaccharides are found in greater abundance in differentiated tissues and are assumed to be playing highly specialised structural roles.

#### Domain 2 : The pectin matrix

The pectin matrix constitutes approximately 30% of the cell wall mass, its chemistry and physical properties are extremely complex and alter throughout development [Roberts 1990] and probably in response to environmental stress. The pectin matrix forms a gelatinous liquid that has often been described as a 'jelly' [Roberts 1990; Carpita & Gibeaut 1993] the physical properties of which, are highly dependent upon the local ratio and concentrations of ions and the local temperature [Walkinshaw & Arnott 1981; Nakajima *et al.* 1981]. The pectin matrix is primarily composed of two highly heterogeneous pectinous compounds, polygalacturonic acids (PGA's) and rhamnogalacturonans (RG's).

PGA's are composed of linear chains of 150-250  $\alpha(1\rightarrow 4)$ -glycosidically-linked sub-units of D-galacturonosyl acid. Hundreds of PGA chains can interact via hydrogen bonding to form helical chains that are about 100 nm in length. PGA's have the unique ability of being able to form, 'junction zones', by coordinating anti-parallel chains of unesterified PGA around calcium ions [Rees 1977; Jarvis 1984;] this behaviour markedly alters the physical properties of the whole gel [Powell *et al.* 1982]. The cross-linking of PGA's via calcium ion co-ordination is highly dependent upon the local concentration of Ca<sup>2+</sup> [Walkinshaw & Arnott 1981], and measurements *in vivo* depict drastic shifts in the ratio and concentrations of magnesium and calcium in plant cell walls during growth and differentiation, these results may indicate development specific roles for PGA's [Nakajima *et al.* 1981].

At a structural level, RG's are contorted rod-like heteropolymers of repeating  $(1\rightarrow 2)\alpha$ -Lrhamnosyl- $(1\rightarrow 4)\alpha$ -D-galacturonosyl acid disaccharide units. RG's are often linked to arabinans, galactans and the highly branched arabinogalactans via the O-4; furthermore, many RG's are acetylated at the O-3 position on galacturonosyl acid sub-units. Many more permutations exist; thus, it isn't surprising to find that as a group RG's comprise some of the most complex polysaccharides yet discovered. Consequently, it has been problematic elucidating both structures and functions for RG's [see Carpita & Gibeaut 1993]. Other pectinous compounds found in the pectin matrix at much lower levels are thought to play significant roles in fundamental cell biology. In particular intriguing evidence has accumulated that a particular sub-type of the polysaccharide grouping, arabinogalactan, plays a crucial role in cell differentiation pathways by acting as a specific cell 'marker' molecule [Pennell & Roberts 1990; Stacey *et al.* 1990].

The pectin matrix in conjunction with the cellulose-xyloglucan framework is proposed to modulate porosity, pH, ion balance and the water content of the plant extracellular environment [Jarvis 1984; Gibeaut & Carpita 1994] and may also play a role to generate signalling molecules [McCann *et al.* 1992]. All of these functions are of course extremely important to the plant, but at a fundamental level, the pectin matrix can even functionally replace the cellulose-xyloglucan framework. When tomato cells are grown in the presence of dichlorobenzonitrile (an inhibitor of cellulose synthesis) the pectin matrix becomes tightly cross-linked to form a semi-rigid cell wall [Shedtetzky *et al.* 1990]. Amazingly, the pectin matrix is effectively acting as a functional replacement for cellulose.

#### 1.2.2 Domain 3 : The plant cell wall proteins

The plant cell wall and the general plant extracellular matrix it occupies play host to a wide range of structural proteins and wall associated enzymes. Each protein group may exhibit substantial inter-species sequence differences, which is especially evident in comparisons between monocot and dicot but all plant cell wall proteins identified to date are glycoproteins. Many of the structural proteins have been reasonably well characterised, as have some of the enzyme constituents. However, many proteins and enzymes associated with the cell wall are as yet still unidentified and of unknown function or significance. Broadly speaking several categories of plant cell wall proteins can be discerned based upon the presence of specific and highly repetitive amino acid motifs. The following is a brief overview of the structure, function and localisation of the known major cell wall structural proteins, table 1.1 presents a collated list of some of characteristics of the plant cell wall proteins considered in the text.

#### 1.2.2.1 Hydroxyproline-rich-glycoproteins

The hydroxyproline-rich-glycoprotein (HRGP) family comprise a highly diverse range of glycoproteins found primarily in the extracellular matrix and predominantly in association with the plant cell wall. As the name suggests all HRGP's are rich in hydroxyproline, however, enough distinctive characteristics have been discovered within the broad

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Table 1.1 – Characterised plant cell wall proteins; collated for this thesis from the references indicated, Somner-Knudsen et al. [1997] and Sachetto et al. [2000].

Type/Group	Species	Repetitive Protein	Protein Structure	Carbohydrate	Localisation	References
		Sequence		composition		
	Soybean	ВРРРКН				
HRGP -	Rape	SPPPVK	Rigid polyproline II-	50-60% (w/w)	Primary cell	Evans <i>et al.</i> [1990]; Memelink <i>et al.</i>
Extensins	Tobacco	ЗРРРРККРҮҮРР	like structure	arabinose & galactose	walls; vascular	[1993]; Smith et al.
	Tomato	SOOOOVKP			lissue	[1980]
HRGP -	Maize					
AGP's	Maize	(Ala-Hyp) <sub>x</sub>	Predicted partial polyproline II-like	>90% (w/w) arabinose & galactose (plus	Vascular bundles;	Kieliszewski <i>et al.</i> [1992]
	Maize	(Ser-Hyp) <sub>x</sub>	structure	some rhamnose & glucuronic acid)	pollen; flowers	
HRGP -	Daturna		Predicted single domain with	~50% (w/w) arabinose	Cell wall:	Desai <i>et al.</i> [1981]; Kieliszewski <i>et al.</i>
SL's	Potato	Ser-(Hyp)4	polyproline II-like structure	& galactose	cytoplasm and vacuole	[1992]
	Arabidopsis thaliana	GGGX				Quigley et al.
GRP's	Carrot	GGX	<ul> <li>Predicted Keratin type structure of</li> </ul>	I	Epidermis;	[1996]; Lin <i>et al.</i> [1996]; Condit &
	Petunia hybridia	GGGX	glycine loops or		vascular tissue	Meagher [1986]
	Atriplex canlsceus	GGGX	allematively p-			
	Soybean	КРРVYK				
PRP's	Maize	КРРТРКР	Predicted B-helices	<3% to >70% (w/w)	Vascular tissue;	Tong <i>et al.</i> [1990]; Stiefel <i>et al.</i> [1988];
	Carrot	РРІНКРР		arabinose & galactose	frequently co-	Chen & Varner
	Rice	РРТҮКР			extensins	[1900], Caeries et al. [1992]

classification of 'HRGP' to distinguish upto 3 individual sub-groupings, the extensins, arabinogalactan proteins and solanaceous lectins.

The extensins are probably the most well known HRGP's and comprise a diverse group of highly basic glycoproteins located in the cell walls of all plants but are particularly prevalent in dicots. They are typically the most abundant protein component, and depending upon the origin of the plant cell wall in question, they may comprise upto 10% of the final dry weight [Cassab & Varner 1988]. Extensins contain upto 40% hydroxyproline, but are also rich in serine, and usually contain some combination of valine, tyrosine, lysine or histidine and a large proportion of the primary amino acid sequence contains the repeating pentapeptide motif, Ser-(Hyp<sub>4</sub>) often within the context of a larger repeating motif [Chen & Varner 1985; Memelink 1988]. The carbohydrate components of extensins usually comprise between 50 and 60% of the total mass. Typically, almost all the hydroxyproline residues within an extensin molecule are glycosylated with mono- to tetra- saccharides of  $(1\rightarrow 4)\beta$ -D-arabinose [Akiyama et al. 1980], furthermore, many serine residues can be  $\alpha$ -linked to a single D-galactopyranose unit [Lamport et al. 1973]. In solution mature extensins conform to a rigid polyproline II type helical structure [van Holst & Varner 1984]. Experimental observations have revealed that mature extensins, (secreted >6 hours previously) are impossible to remove from the cell wall by all non-degradative solvents but newly secreted extensins can be leached from the cell wall with salt solutions [Smith et al. 1984]. These observations do provide direct evidence for the prevailing consensus regarding extensin function, one in which, Carpita & Gibeaut [1993] have likened to the functional equivalent of 'fixing pins' (see figure 1.3), that are inserted into the cell wall to enhance wall rigidity and strength. The method of extensin fixation appears to be achieved via H<sub>2</sub>O<sub>2</sub>-mediated formation of covalent tyrosine cross-links to other extensins, or strong ionic extensin-pectin interactions [Epstein & Lamport 1984].

The second major class of HRGP is the highly soluble, highly acidic and heavily glycosylated arabinogalactan proteins (AGP's). To date all AGP's have been located to the plant extracellular matrix, but precise localisation has been difficult because of their extreme solubility [Fincher *et al.* 1983]. However, some have been found in association with the extracellular side of the plasma-membrane [Pennell *et al.* 1989], and are frequently co-localised with glycine-rich-proteins (see section 1.2.2.3). A typical AGP has a protein moiety that comprises only 2-10% of the total mass, the remainder is a heterogeneous carbohydrate mixture composed of D-galactose and L-arabinose, connected to the protein core via hydroxyproline: $\beta$ -D-galactopyranose linkages [Strahm *et al.* 1981] consequently AGP's can also be classed as proteoglycans [Braam 1999]. The degree of glycosylation observed in AGP molecules ensures that molecular weight estimation is extremely difficult, but all migrate in a pH gradient to exhibit pl's in the

range 2-5. The protein moiety is typically rich in hydroxyproline, serine, alanine, threonine and glycine, and many of the sequenced AGP's exhibit Ala-Hyp repeats towards the N-terminus [Gleeson *et al.* 1989], furthermore, circular dichrometry has revealed that in a similar manner to the extensins a proportion of the protein core (~30%) forms a polyproline II type helix [van Holst & Fincher 1984]. No function has been unequivocally assigned to AGP's, however based on their extracellular location and biochemical and physical properties they have been postulated to act as 'glues' and 'lubricants' [Fincher *et al.* 1983], 'chaperones' of polysaccharide materials during wall remodelling [Gibeaut & Carpita 1994] and cell-cell recognition molecules [Knox *et al.* 1991].

The third class of HRGP, the solanaceous lectins, are extracellular glycoproteins that are structurally similar to extensins and found only in solanaceous plants, the term lectin refers to their ability to bind to carbohydrate materials. Solanaceous lectins can be distinguished from other plant lectins because of their ability to agglutinate oligomers of N-acetyl-β-D-glucosamine [Allen *et al.* 1978; Murray & Northcote 1978], and their unusual amino acid and carbohydrate composition in which hydroxyproline and arabinose are major constituents [Showalter 1993]. Potato tuber lectin is the best characterised of all the solanaceous lectins identified to date. It is a glycoprotein with a molecular weight of 50kDa, and comprises 50% carbohydrate and 50% protein [Matsumoto *et al.* 1983]. The carbohydrate components are primarily tri- and tetra arabinose saccharides linked to hydroxyproline residues, and to a much lesser extent, serine can be linked to individual units of galactose [Ashford *et al.* 1982]. The protein moiety is especially rich in hydroxyproline, serine, glycine and cysteine and in its native state it is exhaustively inter-linked with disulphide bridges so that no free sulphydryl groups are present [Matsumoto *et al.* 1983].

#### 1.2.2.2 Proline-rich-proteins

Proline-rich-proteins (PRP's) are a group of cell wall glycoproteins that have only been identified relatively recently, some members, and perhaps all, contain hydroxyproline, but PRP's can be distinguished from HRGP's because of a much lower degree of proline hydroxylation [Showalter 1993]. PRP's can be sub-categorised into two broad classes that are distinct but not mutually exclusive; those that are normal components of the extracellular environment [Chen & Varner 1985; Datta *et al.* 1989] and those that fall under the special grouping of, 'plant nodulins' [Govers *et al.* 1991]. Both classes can be characterised by a repeating Pro-Pro-Val-X-Lys (X = glutamic acid or tyrosine) motif, which is often incorporated within a variety of other larger repeating units [Hong *et al.* 1990]. PRP's from alfalfa and soybean cell wall's are some of the best characterised to

date and are typified by approximately equal quantities of proline and hydroxyproline [Francisco & Tierney 1990], and are typically only lightly glycosylated [Datta *et al.* 1989].

Developmental-related expression of PRP's has been reported in soybean [Hong *et al.* 1989], potato and tomato [Ye & Varner 1991]. Concurrent data from regulatory studies has indicated extensive PRP involvement in the various stages of development [Jose-Estanyol *et al.* 1992]. PRP's are often found localised at vascular bundles and some can often be found in association with extensins [Wyatt 1992]. PRP's are known to be expressed and oxidatively insolubilised, to form structural wall elements in a similar manner to extensins and glycine-rich-proteins (see section 1.2.2.3) in response to, wounding, endogenous and fungal elicitors, and ethylene treatment [Tierney *et al.* 1988; Marcus *et al.* 1991; Bradley *et al.* 1992].

Plant nodulins are proteins that are specifically produced in plants in response to infection by nitrogen fixing bacteria, and are ultimately constituents of the nodule wall [Franssen *et al.* 1987; Govers *et al.* 1991]. It is proposed that these proteins are involved in nodule morphogenesis and the responses to bacterial infection process in general. Govers *et al.* [1991] hypothesised that plant nodulins may be specifically induced to form an extracellular, 'oxygen barrier' for the  $O_2$ -sensitive nodules.

#### 1.2.2.3 Glycine-rich-proteins

The first glycine-rich-protein (GRP) gene was discovered in 1986 by Condit & Meagher, it encoded an extracellular protein that contains 67% glycine, with the repeating motif Gly-X, where X can be alanine or serine. The Gly-X repeat is a very distinctive characteristic of GRP's and results in a glycine content in the range 25 to 70% [Mundy & Chua 1988]. Secondary structure predictions have indicated that GRP's may exist as  $\beta$ -pleated sheets composed of varying numbers of anti-parallel strands dictated by the particular GRP sequence. Two sub-classes of GRP's exist, the first is developmentally regulated, encodes a signal peptide and is ultimately destined for the exterior of the cell [Keller *et al.* 1988], the second, either lacks a signal sequence or encodes an RNA-binding domain and is cytoplasmic [Sturm 1992].

GRP's are typically the second major structural protein associated with the cell wall of dicot plants [Hirose *et al.* 1994; Carpenter *et al.* 1994; Parsons & Mattoo 1994], though much smaller quantities are also found in monocots [Cretin & Puigdomenech 1990; Didierjean *et al.* 1992]. Commonly they are co-localised with PRP's [Condit *et al.* 1990] and have been frequently located to vascular bundles, particularly the xylem, and are also clearly associated with cells that are going to be lignified [Ye & Varner 1991]. GRP's are commonly expressed during wounding and drought stress, and may provide

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elasticity with a high tensile strength due to extensive  $\beta$ -pleating, or act as nucleation sites for lignification [Keller *et al.* 1988].

#### 1.2.3 The plant cell wall and development

The plant cell wall changes dramatically throughout development therefore it is commonly divided into three sequential stages, cytokinesis, elongation and maturation. At each stage, the chemistry, biochemistry and resultant structure of the cell wall alters.

#### 1.2.3.1 Cytokinesis

After nuclear and cytoplasmic division has taken place the new cells begin to develop a vestigial cell wall called the phragmoplast that physically divides the two cells from one another. The phragmoplast is formed from coalesced Golgi vesicles that primarily contain the polysaccharide, callose [Kakimoto & Shiboaka 1992].

After the initial phragmoplast separation of the two cells, CMF encasing of the cells begins. Deposition of CMFs is closely associated with the terminal ends of the cytoskeleton, thus tentative similarities to animal cell cytoskeletal-extracellular matrix interactions have been proposed [Zhu *et al.* 1993]. Immediately after cytokinesis, CMFs are arranged around the cell in a relatively random fashion (see figure 1.4a). CMFs reorientate as the cell progresses towards elongation, leading to bands of CMFs encasing the cell. This banding can be arranged traversely or longitudinally and ultimately produces a CMF arrangement rather akin to a spring in which CMFs are regularly spaced, presumably by cross-linking to XG polysaccharides [Carpita & Gibeaut 1993] (figure 1.4b).

#### 1.2.3.2 Cell wall elongation

In order for the cell to continue developing it must increase its size by elongating. Crosslinked XGs are considered to be the principal tension bearing components of the cell wall (see figure 1.4b & c), therefore elongation cannot occur until this restraining framework is removed or greatly loosened. Hydrolysis of CMFs and breakage of XG-CMF interactions appears to be the logical process by which the CMF-XG cage could be weakened to allow elongation of the cell. At first glance this appears to be exactly what is happening, pea seedling walls lose up to 50% of their total XG content during elongation [Hayashi 1984]. However, this assumption is directly challenged by recent contradictory evidence presented by McQueen-Mason & Cosgrove [1995]; the walls of cucumber cells that were heat-killed could be induced to elongate, without the detectable release of sugars, by two small, non-hydrolytic proteins, which have been dubbed, 'expansins'. They go on to



#### Figure 1.4

#### The plant cell wall during development.

Information collated and presented in graphical form based on the references quoted.; Terry *et al.* 1981; Epstein & Lamport 1984; Taiz 1984; Tierney *et al.* 1988; Marcus *et al.* 1991; Bradley *et al.* 1992; Pfanz *et al.* 1992; Capita & Gibeaut 1993; McQueen-Mason & Cosgrove 1995. discuss a possible mechanism of expansin disruption of hydrogen bonding between CMFs and XGs.

Thus the process of cell elongation hasn't yet been fully elucidated and other possible mechanisms have been proposed. Most involve interaction with the pectin matrix. For example, PGA's can be de-esterified by the enzyme pectin methyltransferase, thus indirectly altering the local pH. At specific pHs and under the action of auxin, cellulases can hydrolyse XGs *in vitro* [Taiz 1984]. Alternatively, AGP's may bind specifically to XG-CMF cleavage sites thus preventing access by hydrolytic enzymes, slight alterations in the ionic environment as a result of pectin methyltransferase activity could substantially alter the tenacity of AGP's to bind to such sites [Terry *et al.* 1981] (refer to figure 1.4c).

Interestingly, and of particular relevance to this study Fry [1998] and Schweikert *et al.* [2000] have proposed a new theory of cell wall loosening and elongation via the scission of polysaccharides by active oxygen species (AOS), in particular the hydroxyl radical. This theory details a series of Fenton style reactions (see section 1.5 for an overview) that can generate hydroxyl radicals in the extracellular matrix by the action of cell wall and apoplastic peroxidases, providing there is a constant supply of molecular oxygen and a suitable reductant, such as ascorbate. Hydroxyl radicals have been proven to induce polysaccharide scission *in vitro* [Fry 1998; Schweikert *et al.* 2000] a similar situation *in vivo* may ultimately break the restraining XG-CMF framework and consequently loosen the cell wall around the elongating cell. Although hydroxyl radicals are invariable regarded as detrimental, they are so short lived they may act as site-specific oxidants targeted to play useful roles during wall loosening for cell elongation and fruit ripening. The extracellular generation of hydroxyl radicals by this mechanism is covered more fully in section 1.5.2.

#### 1.2.3.3 Cell wall maturation

Once the cell has fully elongated, the cell wall is 'locked' into place. The extensin group of proteins are thought to be important elements of the locking mechanism because it is well established that they are oxidatively fixed into cell wall (see sections 1.2.2.1, 1.4.1 & 1.4.3). Concurrent with this assumption is the observation that cessation of cell elongation, is accompanied by a large increase in the quantity of extensin in the extracellular environment [Sadava *et al.* 1973]. Extensins are insolubilised into the wall by ionic interactions with pectins, and/or covalent linkages via tyrosine bridges to other extensin molecules and PRP's [Epstein & Lamport 1984; Fry 1986] (see figure 1.4e).

In cells that are subsequently going to be lignified GRP and PRP expression is highly colocalised, coupled with this, both protein types rapidly become insoluble after secretion, in a similar manner to extensins [Tierney *et al.* 1988; Marcus *et al.* 1991; Bradley *et al.*  1992]. This possibly indicates that PRP and GRP molecules may interact and are oxidatively insolubilised together. Several potential covalent linkages are feasible between GRP's and PRP's, as is linkage to extensins via isodityrosine linkages (figure 1.4e).

Thus the prevailing hypothesis follows that immediately after cell division CMFs are randomly wound around the cell, and as the cell wall continues to develop the CMFs reorientate to become regularly spaced and transiently fixed into place by XG interaction. During elongation, XG interactions with CMFs are disengaged thus permitting the cell to elongate. As the cell begins to approach maturity, extensins, PRP's and GRP's are expressed and are oxidatively fixed into the wall, thus locking the cell wall into its final conformation around the cell.

#### 1.3 The Apoplast

In 1930 Münch first coined the term 'apoplast' to describe the free space between plant cells through which solutions could move relatively unhindered. He used the term in his description of the theory of translocation of solutes over long distances between sources and sinks within a plant. Since then the term apoplast has been utilised in many ways to describe the extracellular matrix surrounding plant cells. In the 1950's researchers working on extracellular ion movement described the apoplast as [Briggs 1957],

#### "... the free space formed from spaces generated by the continuous wall phase."

Until the late 1950's the term apoplast was used to describe the part of the plant tissue that was essentially, 'non-living' and in which substances in solution could move by diffusion. However during the 1970's plant anatomists further revised the term. They described the apoplast as,

#### "... the walls and intercellular spaces that contain not liquid, but gas".

This definition is distinct from Münch's apoplast and Briggs' free space in that it refers only to the gaseous phase of the intercellular space. During the intervening 20 years the term apoplast has been utilised essentially to refer to the part of the extracellular matrix of plant cells that is distinct from the plant cell wall. Recently Canny [1995] has attempted to clarify and section the apoplast into four distinct types of space with associated putative functions: -

- 1. **The xylem-lumen apoplast:** carries water from roots.
- The water free-space: is composed of the wet intercellular space and is part of the cell-wall space. The liquid effectively forms a film, which permits diffusion among contacting solutions [Altus *et al.* 1985].
- 3. **The Donnan free-space:** a further subsection of the wall space. Contains fixed negative charges.
- 4. **The gaseous intercellular space:** small gas pockets transiently entrapped by water films lining the space.

The apoplast has a multitude of roles to fulfil not least the establishment and maintenance of cell and tissue shape in conjunction with cell walls, turgor development, and uptake and transport of water, ions and metabolites (i.e.: root apoplast and xylem) and gaseous exchange with the external environment. In addition to these many important functions the apoplast is intricately involved in many other cellular reactions and processes, not only under normal growth conditions but also under stress conditions. Abiotic stresses (temperature, ozone, sulphur dioxide, nutrient deficiency or excess, heavy metals, etc) or biotic stress (herbivore and insect induced wounding or bacterial and fungal infections) are known to alter apoplastic functions either as a direct and toxic effect of the stressor or as a reflection of an adaptation to the stress.

The apoplast has an ion and metabolite composition distinct from other cellular compartments, and contains tissue-specific patterns of structural proteins and lytic enzymes. Five distinct processes have been identified that ultimately play roles in defining the metabolic composition of the apoplast [Dietz 1997]:-

- 1. Physiochemical properties (mostly dictated by the cell wall)
- 2. Transport characteristics of neighbouring cells
- 3. Exo- and Endocytotic activity of neighbouring cells
- 4. Water and solute transport in the xylem-lumen apoplast
- 5. Environmental factors (biotic and abiotic)

The apoplast as a homogenous compartment almost certainly does not exist because the factors that contribute towards apoplastic composition are many, varied and some, like external environmental stresses are outside the control of the plant [Canny 1995; Dietz 1997]. Unfortunately there are only a few methods which can distinguish microheterogeneity in the apoplast including; pH measurements using fluorescent dyes [Canny 1995] or microelectrodes [Edwards *et al.* 1988] or immunocytochemistry and tissue prints to investigate protein distribution [Ye & Varner 1991]. Methods in operation to directly sample the apoplast, such as vacuum infiltration, offer low spatial resolution and therefore tend to give average measurements or protein extracts over a relatively large area.

#### 1.3.1 Gaseous exchange

The molecular composition of the apoplastic gas phase is highly dependent upon the composition of the atmosphere, the aperture of the stomata (limits diffusion), the solubilisation profile (gases and salts) of the aqueous phase of the apoplast and the rate of uptake of dissolved material into the cell [Nobel 1983]. The stomata constantly alter aperture size according to the prevailing conditions and thereby restrict equilibrium formation between leaf gas phase and the atmosphere, consequently the water vapour saturation is thought to be approaching 100% in the internal leaf gas space [Farqurhar et al. 1981]. The diffusion resistance for other gases can be inferred from the diffusion resistance to water vapour [Pfanz & Heber 1989]. Laisk et al. [1989] have calculated that under normal atmospheric conditions the concentrations of potential gaseous toxins, such as, ozone, in the leaf gas phase is close to zero. Consequently when elevated atmospheric toxins are present locally, the diffusion gradient is maximal into the gas phase of the leaves, the only limitation to the influx of potential deleterious gaseous toxins is stomatal aperture size. Current theory therefore suggests that the apoplast acts as a very effective sink for gaseous toxins and accordingly the extracellular matrix and the plasma-membrane are effectively exposed targets for immediate damage by such gaseous toxins.

#### 1.3.2 The volume of the leaf apoplast

In all of these studies an important feature for consideration is the relative volumes of gaseous and liquid filled spaces. Using aqueous solutions of low viscosity and known density, measurements have been made of the increase in leaf weight after complete infiltration with a sorbitol solution [Dietz *et al.* 1992; Winter *et al.* 1993 & 1994] and silicone oil [Luwe & Heber 1995]. Taking weight change data, post-infiltration, provides a measure of the volume of intercellular gas space within leaf material. Another way to measure the volume of the apoplast has been to manually analyse ultrathin cross-sections for the relative areas of both intra- and extracellular compartments [Dietz *et al.* 1992; Winter *et al.* 1994]. Such estimates indicate that intercellular gas spaces in leaves typically occupy 18-46% of the total leaf volume and interestingly, the proportion of intercellular gas space to total leaf volume is responsive to stress (i.e.: heavy metal exposure).

The volume of the aqueous phase of the apoplast can be estimated by infiltrating a known concentration of a marker substance; this is effectively diluted by apoplastic

liquid, reducing the concentration of the marker which can then be recovered and assayed to calculate the dilution factor. From this information it is possible to extrapolate the volume of liquid (diluent) in the apoplast [Speer & Kaiser 1991]. Estimates vary and are extremely species-dependent, presumably because of species-specific morphological differences, but approximately 5-20% of leaf water appears to be apoplastic. Table 1.2 presents compiled data detailing the calculated volumes of apoplastic gaseous and aqueous phases for several different plant species.

#### 1.3.3 pH and solute composition of the leaf apoplast

Typically the pH of the apoplast is lower than that of the cytosol, and usually ranges between 5-7. In 1987 Canny used the pH tracer dye sulphorhodamine, and showed that strong acidification (<pH 4) occurs at bundle cells in soybean, i.e.: microheterogeneity in apoplastic pH is evident. Stress, growth substances and enzyme action can all alter apoplastic pH, so one value, for all tissues does not seem likely, more probable is that pH is highly dependent upon local tissue conditions. Changes in apoplastic pH are particularly important because they affect the activity of plant growth regulators such as indoleacetic acid and abscisic acid. Abscisic acid and indolacetic acid respond to altered growth conditions and stresses by redistribution or changes in metabolic activity [Slovik & Hartung 1992]. Table 1.3 illustrates that depending upon the species in question apoplastic pH can range from 5.2 to 7.5 [Dietz 1997].

Mirmura *et al.* [1992] have proposed that the apoplast acts as an ion reservoir for K<sup>+</sup>,  $Ca^{2+}$ ,  $Mg^{2+}$ , Cl<sup>-</sup>,  $NO^{3-}$  and  $PO_4^{3-}$  ions which is ultimately under homeostatic control. Other solutes such as sugars, amino acids, ascorbic acid, dehydroascorbate, phenolic compounds and organic acids can be detected at low but not insignificant levels (see table 1.4). Based on water turnover and solute concentrations detected in the xylem, solutes equivalent to 2 osmol/l<sup>-1</sup> should accumulate in leaves if no solute recycling mechanism was present. Solute cycling between shoot and root has been established for all major elements [Jeschke & Pate 1991]. Using tracer dyes, Smith and Nobel [1986] have observed that solutes tend to accumulate at the extreme tips and margins of leaves; this is especially important for salt-stressed plants, because apoplastic salt-accumulation is thought to be the primary cause of salt-induced necrotic lesions [Speer & Kaiser 1994].

#### 1.3.4 The working definition of the leaf apoplast

Many reports have detailed the use of various types of vacuum infiltration strategies to selectively remove apoplastic components [llett 1998; Hammond-Kosack 1992]. Throughout this project vacuum infiltration has been the method of choice for the

Table 1.2 – Estimates of the volumes of intercellula	ar gas	and	apoplastic	water	spaces
in leaves from different plant species; a	dapted	d fror	n Dietz [199	97].	

Species	Intercellular gas space (μl g <sup>-1</sup> )	Apoplastic water volume (μl g <sup>-1</sup> )	Reference
Carpinus betulus	200	150	Luwe [1994]
Corylus avelana	330	130	Luwe [1994]
Hordeum vulgare	260	53	Winter et al. [1993]
Spinacia oleracea	430	70	Winter et al. [1993]
Spinacia oleracea	450	110	Speer & Kaiser [1991]
Vicia faba	870	110	Hajibagheri et al. [1984]

#### Table 1.3 – Estimates of apoplastic pH in various species; adapted from Dietz [1997].

Species	pH value	Method	Reference
Beta vulgaris	6.0±0.3	Fluorescence	Pfanz & Dietz [1987]
Gossypium hirsutum	6.7-7.5	Pressure chamber	Hartung et al. [1988]
Ruminex lunaria	5.6±0.2	Fluorescence	Pfanz & Dietz [1987]
Spinacia oleracea	6.4±0.3	Fluorescence	Pfanz & Dietz [1987]
Vicia faba	5.2-5.9	Fluorescence	Muhling et al. [1995]

# Table 1.4 – Ion, metabolite and growth regulator composition of the leaf apoplast;adapted from Dietz [1997].

Compound or ion	Apoplastic	Species	Reference
Amino acids	3.1±0.4 mM	Hordeum vulgaris	Lohans <i>et al.</i> [1995]
	3.2±0.8 mM	Spinacia oleracae	Takabama 8 Oniki (1992)
Ascorbic acid	0.45 mM	Spinacia oleracae	
Glucose	1.0±0.4 mM	Hordeum distichum	Tetlow & Farrar [1993]
	4.2±2.3 mM	Hordeum vulgaris	Lohans <i>et al.</i> [1995]
Glutathione	6.5μM	Picea abies	Polle et al. [1990]
Malate	5.5±1.4 mM	Hordeum vulgaris	Lohans <i>et al.</i> [1995]
	0.6±0.3 mM	Spinacia oleracae	
Chloride	4±1 mM	Spinacia oleracae	Speer & Kasier [1991]
	6±2 mM	Pisum sativum	
Magnesium	2±1 mM	Spinacia oleracae	Speer & Kasier [1991]
Phosphate	2.5±0.4 mM	Hordeum vulgare	Mimura <i>et al.</i> [1990]
Potassium	2.4±11.8 mM	Pisum sativum	Long & Widders [1990]
Sodium	1±1 mM	Spinacia oleracae	Speer & Kasier [1991]
	2±1 mM	Pisum sativum	
Abscisic acid	20-100 nM	Gossypium lirsutum	Hartung <i>et al.</i> [1992]
Cytokinin	~10 nM	Gossypium lirsutum	Hartung et al. [1992]
Indoleacetic acid	200 nM	Gossypium lirsutum	Hartung <i>et al.</i> [1992]

selective enrichment of apoplastic materials primarily because of the ease of the technique, the extremely low levels of intracellular contamination and the partial purification achieved during the extraction. As a consequence, with respect to this project, the working definition of the apoplast and derived apoplastic materials, will be defined as, the parts of the extracellular matrix that are recoverable by vacuum infiltration.

#### **1.3.5** Theoretical considerations on the extraction of apoplastic components

Ewald *et al.* [1991] have calculated that the cell walls of higher plants will exhibit sieving properties that will exclude molecules with a Stokes radius larger than 2.9 nm, this corresponds to a molecular mass of 40-45 kDa. Using non-destructive means to extract apoplastic components (i.e.: vacuum infiltration) these sieving effects should be evident. Dietz [1997] reports that a typical whole leaf infiltration eventually subjected to 2-dimensional IEF vs. SDS-PAGE separation will reveal 60-100 major and minor polypeptides with a molecular mass in the range 10-60 kDa. Many of these apoplastic proteins are known to be sub-unit components of larger assemblies with native molecular masses upto 300 kDa [Rohringer *et al.* 1983] for example  $\alpha$ -mannosidase of barley is a complex of 270 kDa [Betz *et al.* 1992]. However, contradictory to the theoretical limits imposed by Elwald *et al.* [1991] apoplastic proteins or polypeptides of this size are readily extractable using vacuum infiltration. There is no obvious explanation to clarify why larger proteins are not sieved out by the cell wall fabric [Betz *et al.* 1992; Dietz 1997].

#### 1.4 Apoplastic Proteins

The secretory system, which comprises the Golgi apparatus, rough and smooth. endoplasmic reticulum and secretory vesicles, delivers secretory proteins and carbohydrate materials (with the notable exceptions of cellulose and callose) to the plant extracellular matrix [see Chrispeels 1991 for a review]. Therefore, the protein composition of the both the plant cell wall and the apoplast is ultimately under the control of the cytosol, this is most obvious when the protein complement rapidly changes in response to biotic or abiotic stresses [Dietz 1997]. The total protein content of the extracellular matrix consists of a covalently bound fraction which acts primarily in structural roles [Fry 1991], some of the key covalently bound proteins have already been described in section 1.2.2. The remainder fall under the umbrella term of apoplastic, as defined in section 1.3.4, and are therefore either ionically bound or freely soluble [Fry 1991] because they are readily removed from the apoplast by vacuum infiltration. Table 1.5 presents some of the better characterised apoplastic proteins. As can be seen all known apoplastic proteins can be broadly categorised into either cell wall, intercellular
Examples of characterised proteins present in the leaf apoplast [compiled and adapted from the references indicated and Dietz 1997 for this thesis]. Table 1.5 –

Functional Grouping	Protein	Characteristics	Specific functions	Reference
	Expansins	Soluble 29 & 20kDa	Cell wall elongation	McQueen-Mason and Cosgrove [1994]
	β-Fructofuranosidase	Soluble and cell wall bound forms; 50-60 kDa	Cleavage of sucrose and regulation of photosynthesis	Sturm & Chrispeels [1990]; Tang <i>et al.</i> [1996]
	Germin/Oxalate oxidase	Soluble homopentameric glycoprotein of 125kDa	The production of hydrogen peroxide via the oxidation of oxalate oxidase; oxidative linkage of cell wall polysaccharides	Lane [1994]
Cell wall related	β-1→3, 1→4-Glucanase	Soluble cell wall maintenance protein	Normal wall metabolism	Hoj & Fincher [1995]
	Hydrolases	Acid pH optima	Breakdown of pectins during cell growth; hydrolysis of ABA glucose esters	Sawicka & Kacperska [1995]
	Lipid transfer proteins	Hydrophilic, ~9 kDa	Cuticle assembly, pathogen-, ABA-, drought-, heavy metal-induced	Molina <i>et al.</i> [1992]]
	α-Mannosidase	270 kDa	Glycosylation and turnover of glycoproteins; may degrade pathogen walls	Betz <i>et al.</i> [1992]; Brune <i>et</i> <i>al.</i> [1994]
	Phosphatase	Two 70 kDa subunits	Hydrolysis of organic phosphates	Ferte <i>et al.</i> [1993]
	Protease	Acidic protein; 70 & 100 kDa	Protein degradation during senescence	Pinedo <i>et al.</i> [1993]
Intercellular	Auxin-binding protein I	Auxin binding domain and KDEL sequence	Putative extracellular auxin binding activity	Brown & Jones [1994]; Shimomura <i>et al.</i> [1999]
signalling	Calmodulin-binding protein	Soluble, apoplastic 21 kDa protein	Putative calmodulin signalling pathway	Jun <i>et al.</i> [1996]
	Chitinase	Multigene family	Hydrolysis of fungal cell walls containing chitin as a component. Defence response	Chen <i>et al.</i> [1994]
	Defensin (formerly $\gamma$ -thionin)	Small 5 Da peptide	Involved in pathogen resistance	Broekaert <i>et al.</i> [1995]
	β-1→3-Glucanase	Hydrolysis of 1→3-β- glucans	Hydrolysis of 1→3-β-glucans present in fungal cell wall. Defence response	Hoj & Fincher [1995]
Stress related	Polygalacturonsidase inhibitor protein	40 kDa	Binds and inactivates fungal polygalacturonsidase	Bergmann et al. [1994]
	Osmotin	25-50 kDa	Drought- and salt-induced	Zhu <i>et al.</i> [1995]
	Peroxidases	Hydrogen peroxide utilising	Cross-linking of cell wall components; degradation of auxin; lignification; roles in defence and stress responses	Pfanz <i>et al.</i> [1992]; Polle <i>et al.</i> [1994]
	Superoxide dismutase	CuZn enzyme	Roles in oxidative stress tolerance	Streller & Wingsle [1994]
	Thionin	Basic, cys-rich, low molecular weight proteins	Inhibition of pathogens	Bohlmann & Apel [1991]

signalling or stress-related with respect to their function. Each will be considered in turn.

## 1.4.1 Cell wall related apoplastic proteins

The plant cell wall is an extremely complex amalgam of polysaccharides and protein materials, (see section 1.2), many cell wall related apoplastic proteins have enzymatic activity which appear to play roles in remodelling cell wall architecture during development.

The cell wall hydrolases are a particular numerous and diverse group of cell wall related apoplastic enzymes that can be found ionically bound to the plant cell wall or freely soluble depending upon the local pH and ionic strength, most characterised apoplastic hydrolases have optimum hydrolytic activity at acidic pH [Hatfield & Nevins 1987; Sawicka & Kacperska 1995]. The grouping is large, and probably contains a repertoire of hydrolases that could potentially break all of the known glycosidic linkages present in the plant cell wall and include examples of cellulases, gluconases, a- & β-galactosidases, βglucuronosidases,  $\alpha$ - &  $\beta$ -mannosidases,  $\beta$ -xylosidases and arabinosidases [Fry 1991]. Some hydrolases, such as the cellulases, were originally considered to play crucial roles hydrolysing the cellulose components of plant cell walls to enable wall loosening and subsequent cell elongation [e.g.: Taiz 1984]. However, this theory has been disputed, most notably by McQueen-Mason & Cosgrove [1995] after the discovery that the small, apoplastic expansin proteins can disrupt CMF-XG interactions, thereby mediating cell wall loosening without recourse to substantial enzymatic digestion of cellulose polymers by cellulases. (Incidentally, Fry [1998] has also proposed a hydroxyl radical theory of plant cell wall loosening described in outline in section 1.2.3.2 and in greater biochemical detail in section 1.5).

Germination, the process of hydration of a dry, mature embryo, that leads to growth of the embryo into a viable seedling involves a huge increase in the mass of the plant, which in wheat escalates from <1 mg to >10 mg with 48 hours, this increase is accompanied by very little cell division but rapid restructuring of the extracellular matrix to accommodate the great increase in cell size [Lane 1994]. Germin is an apoplastic/cell wall associated glycoprotein with a native molecular weight of ~125kDa, that was found to be expressed in tight correlation with the process of germination and cell wall maturation [Lane *et al.* 1992] and was ultimately determined to be an enzyme with oxalate oxidase activity [Lane *et al.* 1993]. Extraction of the protein reveals that there is a tenacious selective association between germin and some members of the arabinogalactan family of extracellular matrix polysaccharides [Lane *et al.* 1987; Jaikaran *et al.* 1990] coupled with the enzymatic production of hydrogen peroxide from oxalate

oxidation the proposed function of germin is the oxidative coupling of polysaccharides into the plant cell wall [Lane 1994].

A range of broad substrate-peroxidases are also proposed to play similar oxidative cell wall strengthening role, and are therefore critical to apoplastic integrity and development. As a group, the apoplastic peroxidases can oxidatively couple polysaccharide-bound phenolic groups including, the *p*-coumaroyl groups of certain polysaccharides and oxidative cross-linking tyrosine residues in extensins and some PRP's and GRP's via isodityrosine bridges to enhance wall rigidity [Epstein & Lamport 1984; Fry 1986]. Furthermore they are integral to lignification via coupling p-hydroxycinnamyl, coniferyl and sinapyl alcohol's [Mader 1980]. Apoplastic peroxidases are often expressed in a tissue-specific, developmental- and stress-related manner; consequently, they are frequently implicated in differentiation, pathogen resistance and wound response [Rao *et al.* 1990; Cordewener *et al.* 1991].

## 1.4.2 Intercellular signalling related apoplastic proteins

The apoplast almost certainly contains intercellular signalling components that individual cells in a multi-cellular organism require to maintain organism integrity. However, unlike animal cells the extracellular signalling systems have not been as well characterised in plants and the literature tends to report tentative similarities to those of animal systems.

In recent years animal systems have illustrated that calmodulin can function as an extracellular signalling molecule by directly binding to and affecting the activity of other proteins. As an example Inui *et al.* [1990], have shown that calmodulin can actively prevent the binding of thyroid hormone to its target membrane receptor, thereby altering the cellular response to the hormone. Apoplastic calmodulin was first reported by Biro *et al.* [1984] and has since been verified by a number of different workers [Ye *et al.* 1989; Sun *et al.* 1995]. The exact role that calmodulin plays in plant intercellular signalling and control mechanisms is not clear but Sanchez *et al.* [1989] have reported that calmodulin can bind to and inactivate a cell wall peroxidase from *Cicer arietinum*, this evidence may point to an extracellular role for calmodulin in plant cell development, as has been verified in animal systems [MacNeil *et al.* 1988]. Recently, Jun *et al.* [1996] has reported the isolation of a 21 kDa apoplastic calmodulin binding protein and proposes a tentative Ca<sup>2+</sup>:calmodulin based signalling system similar to the known intracellular system [Trewavas 1991; Jun *et al.* 1996].

The actions of the plant growth substance, auxin are reasonably well characterised after years of research. On the whole plant scale the hormone modulates root and shoot formation, elongation and growth, and at the molecular level induces membrane

hyperpolarisation and acidification of the extracellular matrix. The latter effects may go someway to explaining how auxin can induce cell elongation and growth in auxinsensitive tissue [see Millner 1995 and Palme & Gälweiler 1999 for reviews]. Despite the efforts of a multitude of researchers, to date, there is no concrete evidence of any of the components of the auxin signal transduction pathway. There have, however, been a number of putative auxin-receptors reported, the most notable and best characterised is auxin-binding protein (ABP) 1 which was isolated from maize in 1985 [Löbler & Klämbt]. ABP1 is a dimer of 40 kDa [Löbler & Klämbt 1985] that contains an auxin-binding motif, a glycosylation site and a carboxy-terminal KDEL signal sequence that targets and retains the protein in the endoplasmic reticulum [Campos et al. 1994]. The KDEL targetting sequence is difficult to reconcile with the current consensus on the site of auxin perception, the plasma-membrane [Venis & Napier 1995; Millner 1995] and additionally both standard immunofluorescence microscopy [Napier et al. 1992; Brown & Jones 1994] and density gradient fractionation data [Shimomura et al. 1988] further indicate that the vast majority of ABP1 is located to the endoplasmic reticulum. However, recently, the use of highly sensitive immunogold-label electron microscopy [Jones & Herman 1993] and silver-enhanced immunofluorescent microscopy [Deikmann et al. 1995] has visualised a small sub-population of ABP1 present in the extracellular matrix, in the cell wall and clustered at the outer surface of the plasma-membrane. It is therefore probable that this small sub-population of ABP1 is functionally resident in the apoplast [Brown & Jones 1994], and although unidentified at the present, the ABP1-auxin complex is proposed to interact with a plasma-membrane receptor, thus mediating the auxin signal to the cell.

In addition to the obvious role of ABP's in trafficking auxin from the apoplast to the plasma-membrane to induce cellular changes, auxin is also likely to interact with the proteins involved in auxin metabolism and extracellular and apoplastic auxin related effects [Cohen & Bandurski 1982; MacDonald *et al.* 1991]. In support of this hypothesis photoaffinity labelling has revealed a number of enzymes that are capable of binding auxin, these include, a hydrolase [Brzobohaty *et al.* 1993], a 1,3- $\beta$ -glucanase [MacDonald *et al.* 1991], a glutathione-S-transferase [Bilang *et al.* 1993; Reinard & Jacobsen 1995] and a superoxide dismutase [Feldwisch *et al.* 1994].

## 1.4.3 Stress related apoplastic proteins

Any stress whether abiotic or biotic disturbs the normal physiological functioning of any organism. Typically, stress responses are transmitted primarily from the cells to the apoplast, but in certain cases, such as pathogen invasion and ozone stress, mediators are generated or liberated in the apoplast and transmitted to the cell. Plants, like other organisms, have developed appropriate responses to deal with stresses that ultimately

serve to re-establish normal physiology and in effect remove or reduce the stress. A common adaptation to a variety of stresses is the alteration of the extracellular matrix; this includes physical restructuring and alteration of the biochemical properties of the plant cell wall, modulation of the solute composition of the apoplast and changes to the apoplastic protein complement. Drought, excessive salinity, heavy metals, excessive cold, air pollutants, pathogens, and other stressors have all been shown to modify both qualitatively and quantitatively the protein composition of the apoplast. Salt, heavy metal stress, pathogen invasion, ozone, elevated SO<sub>2</sub> [Pfanz *et al.* 1989] and cold-hardening [Marentes *et al.* 1993] all produce increases in apoplastic protein levels, whereas, drought reduces the quantity of extractable apoplastic protein. Stress can be divided into two groupings, biotic, which involves attack from a biological agent and in which the plant has a specific set of defensive reactions (see section 1.4.3.2 & 1.5.2) and abiotic, primarily stemming from a local environmental change that instigates a variety of adaptations to ensure survival (section 1.4.3.1).

## 1.4.3.1 Abiotic stress

## Temperature

Chilling-sensitivity (6-4°C) in plants arises primarily because of two factors; firstly, at a critical temperature, lipids present in the membrane phase shift from a liquid-crystalline state to a solid state, this critical temperature is determined by the ratio of unsaturated to saturated fatty acids in the membrane [Quinn 1988]. Many chilling- and frost-tolerant plants can increase the relative quantity of unsaturated fatty acids, thus permitting membranes to remain fluid and therefore functional at lower temperatures. The second major factor is the inadvertent generation of excessive quantities of active oxygen species; this will be discussed further in section 1.5.2.

Frost-tolerance in plants enables survival below freezing temperatures; thus, additional protective mechanisms other than adjustment to membrane fatty acid composition must be active. Frost-tolerant plant cells are known to dehydrate by losing water to small ice crystals that form in their apoplast. Recently, Marentes *et al.* [1993] and Griffith *et al.* [1992] have reported the isolation of apoplastic antifreeze, and ice-nucleation proteins from rye. The Puma and Musketeer rye varieties can survive temperatures as low as – 29°C [Fowler *et al.* 1977] because as the temperature drops, the ice-nucleation and antifreeze proteins are secreted into the apoplast. The ice-nucleation proteins promote the formation of many ice crystals while the antifreeze proteins of both apoplastic proteins ensure that any ice crystals that do form are very small and numerous, consequently these proteins prevent physical damage to the cells from unstrained ice

crystal formation. The roles these proteins play in controlling ice crystal formation are crucial for the plant to survive sub-zero temperatures.

All organisms respond to rapid increases in environmental temperature by transcriptional activation of genes that encode heat shock proteins. Heat shock proteins function to protect the cell from the damaging effects of over-heating by: -

- 1. Switching gene expression programmes [Craig et al. 1994].
- 2. Binding to incorrectly folded proteins and refolding them into the correct conformation [Jacob & Buchner 1994].
- 3. Aiding in compartment targetting and secretion [Parsell & Lindquist 1993].

No evidence appears to have been proposed to suggest how elevated temperatures may affect apoplastic components, though, the stress responses of plants to prolonged periods of elevated temperatures may be similar to that observed in drought, because of increased water loss (see below).

## Drought

Plant growth is immediately affected by sub-optimal water supply, this effect is not due to lowered turgor but seems to stem from alterations to the physical properties of cell walls [Zhu & Boyer 1992]. No consensus appears to have been reached detailing the exact changes to cell walls during drought stress. However, a temporary cessation of leaf elongation, for approximately 30 minutes, was observed when roots were exposed to a hyperosmotic solution of mannitol. Leaf elongation quickly recovered because of changes to the elastic properties of the cell wall [Serpe & Matthews 1992] the exact molecular basis for such rapid modulation of cell wall structural properties is unknown. Changes to the apoplastic protein complement have been observed during drought. Covarrubias *et al.* [1995] extracted two basic polypeptides of 33 and 36 kDa from bean seedlings, which are cell wall associated, induced, and secreted into the apoplast in response to drought stress. Unfortunately, no function has yet been attributed to either polypeptide.

One important component of the drought stress response is undoubtedly abscisic acid. Abscisic acid induces stomatal closure and promotes genetic adaptation to drought [Bray 1990]. Computer simulations modelling drought stress show that abscisic acid would be redistributed in a pH-dependent fashion, with a particular increase in the levels of apoplastic abscisic acid. In turn, elevated levels of apoplastic abscisic acid are known to trigger stomatal closure [Slovik *et al.* 1995] that ultimately reduces water loss and consequently lessens drought stress.

## Salt

An elevated level of salt at the roots results in an accumulation of ions in the shoot apoplast of sensitive species. This accumulation of ions produces the common symptoms of salt-stress, wilting and necrotic lesions on leaves, presumably due to altered osmotic and turgor properties [Niu *et al.* 1995]. These symptoms may be due to the deposition of excess chloride ions in the apoplast, which in turn, would affect the osmotic potentials and the ionic properties of the plant cell wall; 18-day old pea (*Pisum sativum*) seedlings were watered with a 50mM NaCl solution and for the next four days the concentration of apoplastic solutes from leaves remained relatively stable. After day 4 apoplastic chloride ion concentrations of a range of other ions were also increased by a factor of 10 [Speer & Kaiser 1994]. The authors speculate that this general accumulation of ions in the apoplast results from a non-specific release of intracellular solutes.

More rapid and drastic changes occur to the cell wall and apoplastic protein complement during salt-stress situations. Bressan *et al.* [1990] investigated these effects on tobacco cell cultures. Salt-adapted cells had walls containing, 50% less polysaccharides, 40% less pectins, three times as much insoluble protein. Exactly how this affects plant cell wall structure has not yet been determined, but, a similar study with the algae, *Nitella*, has revealed that more than 50% of the walls' cation exchange capacity was completely lost [Gillet *et al.* 1992]. Salt-adapted tobacco cells also secreted seven times more protein into the medium, similarly, apoplastic extracts from the intact leaves of salt-stressed tobacco plants exhibit quantitative and qualitative changes in protein complement [Ramanjulu & Dietz, unpublished data]. One of these proteins is probably osmotin [Singh *et al.* 1987]. Osmotin and osmotin-like proteins are induced by abscisic acid, NaCl, wounding, ethylene, viral infection and fungal pathogens, their exact role in salt-stress responses is undetermined [Zhu *et al.* 1995].

## Heavy metals

At elevated levels, Cd, Ni, Zn and Al all affect plant growth [Woolhouse 1983; MacNair 1993]. All heavy metals eventually accumulate in the root, shoot, and most notably in the leaf apoplast and the vacuole, unless the plant has specific excretion mechanisms [Brune & Dietz 1995]. The apoplast and vacuole appear to act as an effective, 'repository' to ensure that metal ions do not accumulate in the cytoplasm at levels that would cause cellular disruption and ultimately cell death.

Accumulation of metal ions in the apoplast alters the physical and chemical properties of the cell wall. The concentration of metal ions in the apoplast dictates whether the ions will displace Ca<sup>2+</sup> from anionic sites in the cell wall. The ionic cross-linking that exists in the cell wall is extremely important to normal wall structure and function. This mechanism has been proposed as the primary cause of the Al-dependent reduction of root elongation observed at sub-toxic concentrations [Jones 1995].

Heavy metals appear to induce more quantitative than qualitative changes in the complement of apoplastic proteins. 1 and 2-dimensional patterns of heavy metal stressed apoplastic extracts, reveal enhanced levels of proteins in the 15-30 kDa range. It is possible that heavy metal stress may induce an elevated secretion of all apoplastic proteins in this molecular weight region and <sup>35</sup>S-methionine labelling and assay of apoplastic protein does appear to confirm elevated production and secretion of proteins into the apoplast [Brune *et al.* 1994; Blinda *et al.* 1997]. However, Blinda *et al.* [1997] have noted that a more probable reason for apparently elevated levels of apoplastic proteins is that the altered properties of the cell wall, induced by exposure to the heavy metal ions, would probably permit the easier extraction of associated proteins.

## 1.4.3.2 Biotic stress

Bostock & Stermer [1989] define a wound as;

## "... an internal or external injury that breaks the outer protective layer of a plant, that ultimately leads to cell destruction in a specific area of tissue."

Plants can be wounded by a variety of abiotic (wind, rain, hail, freezing) and biotic (abrasion during growth, herbivores, pathogens) means. Consequently, it is not surprising that Jongsma *et al.* [1994] and Baron & Zambryski [1995] have speculated that only one signal transduction pathway produces the wound response, irrespective of the means of wounding. This wound-defence response is exclusively induced by 'elicitor' compounds, which are present, either as a result of localised physical damage, (cell wall oligosaccharides and other wall components [Bishop *et al.* 1984; Darvill & Albersheim 1984; Lamb *et al.* 1989]) or, originate from an invasive pathogen (glycans, oxalic acid, salicylic acid or hydrolytic enzymes [Lamb *et al.* 1989; Davis *et al.* 1992; Bi *et al.* 1995; Ebel & Cosio 1994]). Examples of plasma-membrane receptors for both microbial and plant cell wall derived elicitors have been isolated [Apostol *et al.* 1989; Horn *et al.* 1989]. Additionally, there is also evidence that certain AOS may also act as 'secondary elicitors', the strongest evidence so far points to hydrogen peroxide produced during the oxidative burst [Tenhaken *et al.* 1995] (see below).

The initial rapid defence response has been suitably called the 'oxidative burst', because large quantities of active oxygen species (AOS) are liberated in the extracellular matrix [Wojtaszek 1997] (the biochemical side of this topic is dealt with in greater detail in section 1.5 but see figure 1.5 for a diagrammatic overview). AOS's are toxic intermediates of successive reductions of molecular oxygen, generating, superoxide anions, hydrogen peroxide and the hydroxyl radical [reviewed in Mehdy 1994]. The hydroxyl radical is extremely reactive and can cause lipid peroxidation, enzyme inactivation and degradation of nucleic acids, consequently it is a very effective anti-microbial agent. Typically, AOS's are the first response to a wound or potential infection, for instance, in soybean suspension cultures  $H_2O_2$  in the medium is detected after <5 minutes of exposure to an elicitor [Apostol et al. 1989]. The oxidative burst generates many of the characteristics of the early response. For instance, 2-5 minutes after the production of  $H_2O_2$  has begun, existing proline- and tyrosine-rich proteins are oxidatively insolubilised into the cell wall [Bradley et al. 1992; Brisson et al. 1994] and localised programmed cell death of pathogen infected cells begins to occur [Levine et al. 1994]. Furthermore, Matern et al. [1995] have reported the rapid deposition of wall strengthening, lignin-like and phenylpropanoid materials in the apoplast in and around the infection site that once deposited greatly reduce the digestibility of the tissue. On a gross scale all of these early responses ultimately deprive the pathogen of nutrients thereby limiting its potential to establish an infection and spread.

Elicitors are potent inducers of defence gene expression, for example, the elicitor, hepta- $\beta$ -glucoside, can induce phytoalexin synthesis in soybean at only 1x10<sup>-8</sup> M concentration [Albersheim *et al.* 1992]. Defence-related genes are induced both locally and distally in response to the infection process [Stintzi *et al.* 1993], this requires the activation of gene expression programmes and the transcription and translation of new mRNAs, thus these responses typically become apparent hours after the initial insult [Brisson *et al.* 1994].

General defence-related proteins such as, serine proteinase inhibitors [Graham *et al.* 1986], prosystemin [McGurl *et al.* 1992], polyphenol oxidase [Constabel & Brisson 1995], leucine aminopeptidase and cysteine proteinase inhibitors [Hildmann *et al.* 1992] are synthesised and secreted into the apoplast. In additon, specific pathogenesis-related proteins are also synthesised in response to pathogen infection. In general, these proteins are able to inhibit the growth and the spread of the pathogen and are resistant to the harsh conditions in and around the infection site (acidic pH and proteolytic cleavage by pathogen enzymes) [Niderman *et al.* 1995]. Pathogenesis-related proteins can be classified into several categories, lytic enzymes such as, chitinases and  $\beta$ -1,3-glucanases [Fleming *et al.* 1991; Rasmussen *et al.* 1992; Hoj & Fincher 1995] which digest microbial cell walls, ribosome-inactivating proteins [Leah *et al.* 1991], chitin-binding proteins [Hejgaard *et al.* 1992] and thaumatin-like proteins [Vigers *et al.* 1991].



Figure 1.5

# Events leading to the production of active oxygen species (AOS) and nitric oxide (NO).

Figure prepared for this thesis; refer to the text for a more detailed explanation of the figure. Solid arrows indicate identified pathways and events. Broken arrows indicate probable pathways and events. Key: E=elicitor; R=receptor; cNOS=constituitively expressed nitric oxide synthase; NO=nitric oxide; O<sub>2</sub>=molecular oxygen; H<sub>2</sub>O<sub>2</sub>=hydrogen peroxide; H<sub>2</sub>O=water; GSSG=glutathione; GSNO=nitroso-glutathione; O<sub>2</sub>=superoxide anion; OH =hydroxyl radical; OONO =peroxynitrite radical; O2H=hydroperoxyl radical; SOD=superoxide dismutase; CAT=catalase. However, other anti-microbial compounds are also produced in response to pathogen attack. Phytoalexins and the  $\alpha$ -,  $\beta$ - and,  $\gamma$ -thionins [Bohlmann 1991] are a range of low molecular weight anti-microbial compounds with a diverse range of action. The thionins are extremely important in most plant species, and are thought to act by binding to ion channels, thus throwing microbial signalling into disarray.

## 1.5 Active Oxygen Species & Nitric Oxide

Active oxygen species (AOS) as a group comprise the superoxide anion ( $O_2$ ), hydrogen peroxide ( $H_2O_2$ ) and the hydroxyl radical (OH), all of which are potentially dangerous to macromolecules because of their strong oxidising properties; the redox chemistry of AOS is detailed in section 1.5.1. Even though AOS are potentially deleterious, there are mechanisms in place in plants to purposefully produce appreciable quantities of AOS during defence responses and development, and many of the normal processes of life generate AOS as by-products; these are discussed in section 1.5.2. However, due to their inherent toxicity, plants have developed an array of anti-AOS mechanisms to 'mop up' any excess in order to prevent cellular damage, however, in times of stress these defences may be overcome. Section 1.5.3 illustrates that this phenomenon is common to many types of stress that ultimately perturb the redox state of cells. Recently, evidence has been accumulated that the radical nitric oxide (NO) is also implicated in active oxygen metabolism and signalling, these ideas will be explored in section 1.5.4.

## 1.5.1 The redox chemistry of active oxygen species

The generation of active oxygen species in biological systems is not an unexpected phenomenon. The oxygen molecule itself is a di-radical, because it contains two electrons that reside in different orbitals, these electrons are not spin paired, therefore, thermodynamically, two more electrons are required to fill the orbitals. However, the oxygen molecule cannot accept spin matched pairs of electrons until one of its unpaired electrons undergoes spin reversal; at ambient pressure and temperature the collision period between molecules is so short that spin reversal faces a large kinetic barrier. Enzymatic oxidation systems effectively remove this kinetic barrier because the collision period between oxygen and the substrate is lengthened at the active site producing a controlled oxidation of the substrate molecule [reviewed in McCord 2000]. However, in biological systems one electron transfers are common (eg: cytochrome chains or photosynthesis) and consequently it is relatively easy for molecular oxygen to undergo a one step reduction and form the superoxide anion ( $O_2$ ), this molecule rapidly establishes an equilibrium with a proton to form the superoxide anion-hydroperoxyl radical ( $O_2$ H) acid-base pair:-

The half-life of the superoxide anion-hydroperoxyl radical acid-base pair is less than 1 second at neutral pH and rapidly shortens as the pH becomes more acidic [Sutherland 1991]. The pair spontaneously dismutates into another AOS, hydrogen peroxide ( $H_2O_2$ ):-

$$2 H^{+} + 2 O_{2}^{-} \rightarrow H_{2}O_{2} + O_{2}$$
 (3)

However, while the superoxide anion does exist it can also act as a reducing agent for transition metals such as,  $Fe^{3+}$  and  $Cu^{2+}$ :-

The generation of the reduced forms of copper and iron opens up the possibility of Fenton style reactions to generate the most reactive and deleterious of all AOS, the hydroxyl radical (OH):-

These latter reactions are particularly avid, for instance, even metal ions that are complexed within proteins or chelators are capable of generating the hydroxyl radical from hydrogen peroxide-mediated oxidation [Mehdy 1994].

## 1.5.2 The biological production of active oxygen species

Figure 1.6 diagrammatically presents the major known sources of AOS in plants. Two important intracellular sites of constant AOS production are the mitochondria and chloroplasts. Mitochondria typically convert 1-2% of the total molecular oxygen used for respiration into the superoxide anion, specifically, complex I, partially reduced ubiquinone and cytochrome *b* in complex III are all energetically favourable points within the electron transfer chain that single electron reduction of molecular oxygen can readily occur [Richter *et al.* 1995; McCord 2000]. It has been calculated that a single mitochondrion, operating normally would generate 3x10<sup>7</sup> superoxide anions per day [Richter *et al.* 1995]. Similarly, the involvement of molecular oxygen in photosynthesis exposes the photosynthetic apparatus to the inadvertent production of AOS. In fact, the single electron reduction of oxygen to form the superoxide anion is known to occur primarily in photosystem I [Asada *et al.* 1974] and more recently discovered in photosystem II during strong illumination [Klimov *et al.* 1993; Ananyev *et al.* 1994].

The production of AOS during plant-pathogen interaction has been briefly covered in section 1.4.3.2 and figure 1.5 diagrammatically illustrates the current consensus



## Figure 1.6

## Sources of active oxygen species in plants

Figure prepared for this thesis; refer to section 1.5.2 for detail. Active oxygen species are generated at five known sites in plants; mitochondria, chloroplasts, by a putative plasmamembrane NADPH-oxidase complex [Sutherland 1991; Vera-Estrella *et al.* 1992], and by apoplastic and cell wall derived peroxidases [Fry 1998]. Key- O<sub>2</sub>=superoxide anion; OH=hydroxyl radical.

mechanism. In brief, the interaction of the pathogen with the extracellular matrix leads to the release of elicitor compounds [Bishop *et al.* 1984; Darvill & Albersheim 1984; Lamb *et al.* 1989; Bi *et al.* 1995; Ebel & Cosio 1994] which bind to and activate plasmamembrane receptors [Apostol *et al.* 1989; Horn *et al.* 1989], triggering the G-protein mediated opening of ion channels that induce hyperpolarisation of the plasmamembrane and the influx of calcium ions. Elevated cytosolic calcium may activate a nitric oxide synthase enzyme (see section 1.5.4) and a putative plasma-membrane-associated NADPH-dependent oxidase complex [Sutherland 1991; Vera-Estrella *et al.* 1992]. The latter transfers single electrons from the NADPH molecule to molecular oxygen thereby generating and liberating the superoxide anion into the apoplast. Once in the apoplast the superoxide anion either undergoes spontaneous (see section 1.5.1) or enzymatic dismutation (see figure 1.5, table 1.6 and section 1.5.3), to hydrogen peroxide, as a consequence, hydrogen peroxide is the major AOS observed during the defence response to pathogens.

The hydroxyl radical generated by the Fenton reaction (see section 1.5.1) is known to be able to decompose cellulose-containing materials such as straw and sawdust [Halliwell 1965]. Miller [1986] illustrated that certain cell wall polysaccharides could undergo mild non-enzymatic scission when incubated with hydrogen peroxide. Recently, Fry [1998] has extended these observations to prove that hydroxyl mediated scission of polysaccharides is possible in simple reactions mixtures that deliberately resemble the apoplast; the basic requirements are molecular oxygen, a transition metal such as Cu<sup>2+</sup>, an appropriate electron donor such as ascorbate and a source of hydrogen peroxide - all are found in the apoplast. The hydroxyl radical is proposed to be formed via the Fenton reaction (reaction scheme 7, section 1.5.1) and the reduced form of the copper ion from the redox reaction with ascorbate:-

## ascorbate + $Cu^{2+} \rightarrow Cu^{+}$ + dehydroascorbate

However, the entire mechanism pre-supposes a ready supply of hydrogen peroxide. Germin (an enzyme that readily generates large quantities of hydrogen peroxide during germination in barley) [Lane 1994], some peroxidases when provided with a suitable reductant [Wojataszek 1997] and the putative plasma-membrane NADPH-oxidase complex [Sutherland 1991; Vera-Estrella *et al.* 1992; Fry 1998] can all generate substantial quantities of hydrogen peroxide. The hydroxyl radical appears to cause scission of the glycosidic linkages in the polysaccharide 'backbone' [Schuchmann & von Sonntag 1978]; this has prompted Fry [1998] and Schweikert *et al.* [2000] to propose that this is a potential mechanism of cell wall loosening during cell elongation (see section 1.2.3.2).

## 1.5.3 Anti-AOS defences & oxidative stress

The mitochondria and chloroplasts are constantly exposed to constitutively generated AOS, consequently, both organelles are especially rich in molecules capable of scavenging AOS such as, ascorbate, vitamin E, glutathione and ubiquinol-10 [Chance *et al* 1979; Richter 1995] and contain high levels of superoxide dismutases, peroxidases and catalases for the enzymatic breakdown of AOS (see table 1.6 for mechanisms of action). Under physiologically ambient conditions, these mechanisms for the management of AOS are sufficient to ensure that no overt cellular damage arises from AOS generated by either organelle. However, under stress these mechanisms can be swamped and may lead to macromolecular oxidation, cellular injury and ultimately the symptoms of stress [Purvis *et al.* 1995; O'Kane *et al.* 1996].

AOS are specifically generated in the apoplast during the biotic stress response and at specific stages of development (see section 1.5.2 and figure 1.5). The apoplast does contain a range of anti-AOS defences, such as, ascorbate, phenolic substances and glutathione [Polle *et al.* 1990; Takahama *et al.* 1992; Luwe *et al.* 1995] which are effective scavengers of AOS, and anti-AOS enzymes such as, superoxide dismutases, catalase, ascorbate peroxidase and glutathione reductase enzymes [Edwards & Dixon 1991; Streller & Wingsle 1994; Vanacker *et al.* 1999], that are capable of catabolising AOS. The actions of these molecules and enzymes in the apoplast is rather low when compared to whole leaf extracts, but their concentration in this small compartment with such a low water volume per gram of tissue (see table 1.2) is actually relatively high [Streller & Wingsle 1994; Vanacker *et al.* 1999].

These apoplastic anti-AOS mechanisms are responsive to oxidative stress generated from the external environment, biotic attack and development. As an example, atmospheric ozone concentrations have steadily increased over the past 50 years [Chameides *et al.* 1994] and elevated levels can cause foliar injury, reduce shoot and root growth, induce premature senescence, reduce photosynthetic capacity [Heath 1994] and produce increased susceptibility to infection. Ozone enters the plant via the stomata (see section 1.3.1), and is rapidly dissolved into the apoplastic liquid where it readily degrades into several AOS, most notably the hydroxyl radical (OH<sup>-</sup>) which can oxidise most chemical and biological molecules. Luwe *et al.* [1993] have demonstrated that apoplastic levels of ascorbate fall from 420 $\mu$ M to 50 $\mu$ M within 6 hours of treatment with 640 $\mu$ g m<sup>-3</sup> ozone (very high levels) in spinach leaves. Luwe & Heber [1995] later

## Table 1.6 – Characterised anti-AOS defences; compiled from the references indicated for this thesis.

Type of anti-	Name	Mode of action	Location	References
AOS				
	Superoxide		Apoplast and	Strollor 8
Enzymatic	dismutases	$2^{\cdot}O_2^{-} + 2H^{+} \rightarrow H_2O_2 + O_2$	intracellular	Streller & Wingsle [1994]
			locations.	
	Peroxidases	compound $+ H_{-}O_{-} \rightarrow$	Apoplast and	Willekens <i>et al.</i>
		redcompound + H <sub>2</sub> O <sub>2</sub> → oxcompound + H <sub>2</sub> O	intracellular	[1995]; Takahama &
			locations.	Oniki [1992]
	Catalases	$2 \; \mathbf{H_2O_2} \rightarrow \mathbf{H_2O} + \mathbf{O_2}$	Apoplast and	Willekens et al.
			intracellular	[1990]
			locations.	-
	Ascorbate	eccerbate (red)	Apoplast and	Luwe [1996];
		dehydroascorbate (ox)	intracellular	[1990];
		· · ·	locations.	
Scavenging	Glutathione	<sub>red</sub> glutathione → <sub>ox</sub> glutathione	Apoplast and	Luwe [1996];
			intracellular	[1998]
			locations.	
	Polysaccharides			Fry [1998];
		of carbohydrate chains	Apoplast	al. [2000]
	Dhanallas			
	Phenolics	r <sub>ed</sub> phenolic compound →	Anonlast	Takahama &
		<sub>ox</sub> phenolic compound		Oniki [1992]
	Ubiquinol			Arrovo ot cl
		r <sub>red</sub> ubiquinol → <sub>ox</sub> ubiquinol	Membranes	[1998]
	I		· · · · · · · · · · · · · · · · · · ·	<u> </u>

Note - AOS are indicated in bold type; key; red=reduced; ox=oxidised.

conducted a series of experiments at ozone levels more representative of those found in certain ozone 'hotspots' in the atmosphere and no significant alterations in apoplastic ascorbate redox state were found, but the leaves still eventually developed the symptoms of ozone toxicity. One explanation of this seemingly contradictory data are that the localised depletion of ascorbate in and around the apoplast of the sub-stomatal cavity did occur and therefore contributed to cell damage and ultimately chlorosis but because infiltration techniques, with the inherent problem of low spatial resolution, were used to sample the apoplast, it is unlikely that the authors would be able to detect changes in very small areas.

Anyhow, the assumption that apoplastic ascorbate is crucial to cellular protection from AOS pre-supposes that mechanisms exist to continually replenish apoplastic stocks. In fact, Horemans et al. [1994] and Rautenkranz et al. [1994] have reported that dehydroascorbate, the oxidised product of ascorbate, is continually being recycled by a reductive mechanism, when tissues are exposed to sub-toxic levels of ozone. Two potential mechanisms of ascorbate recycling have therefore proposed and are diagrammatically presented in figure 1.7. The first, the ascorbate exchange theory (figure 1.7 [A]) proposes that apoplastic ascorbate could be maintained in a reduced state by continual replenishment from neighbouring cells, thus, the redox state of the apoplast would be directly coupled to that of the local cytoplasm. In support Rautenkranz et al. [1994] have reported that barley protoplasts are able to transport ascorbate and dehydroascorbate across their plasma-membranes, and perhaps tellingly, the rate of transfer into the cytoplasm is approximately twice as fast for dehydroascorbate when compared to ascorbate. The alternative mechanism rests with the fact that the plasmamembrane of plant cells contains a cytochrome b complex (figure 1.7 [B]). Horemans et al. [1994] have inserted this complex into the membranes of vesicles containing ascorbate. When dehydroascorbate is added externally, the ascorbate solution inside the vesicle is rapidly reduced to dehydroascorbate. A similar mechanism may be in action at the plasma-membrane to replenish apoplastic ascorbate in vivo.

## 1.5.4 The chemistry and the roles of nitric oxide

In 1992 the molecule, nitric oxide (NO) was recognised by Science magazine as, 'Molecule of the Year', because of its fundamental and wide ranging effects in animal systems [Wink 1998]. Until relatively recently the effects of NO in plants had primarily focused upon NO as an atmospheric pollutant, its uptake into leaves and subsequent metabolism and toxicity [Hufton *et al.* 1996]. However, the production of biologically active NO in plants is undisputed [Durner & Klessig 1999] but the exact mode of NO production is still under debate. In animal systems NO is exclusively generated by the enzymatic oxidation of L-arginine: -



## Figure 1.7

# Ascorbate recycling theories.

Figure prepared for this thesis; see section 1.5.3 for detail. [A] = diagrammatic representation of the ascorbate exchange theory [Rautenkranz et al. 1994]; [B]

= diagrammatic representation of the ascorbate regeneration theory [Assard et al. 1992; Horemans et al. 1994].

L-arginine +  $O_2 \rightarrow$  L-citrulline + NO (1)

The reaction is catalysed by the enzyme nitric oxide synthase with the aid of NADPH, flavin, thiol and tetrahydrobiopterin cofactors [Poulos & Raman 1998]. Similar enzyme activity has been reported in plants [Cueto *et al.* 1996; Ninnemann & Maier 1996] but neither the protein nor the gene encoding the protein has been identified [Wojtaszek 2000] furthermore, the co-factor tetrahydrobiopterin is currently not known to be synthesised in plants [Durner & Klessig 1999]. However, plants are already known to contain a range of enzymes with the potential to generate NO, the NAD(P)H-dependent nitrate or nitrite reductases, therefore these enzymes have also been proposed to produce the bulk of NO in plants [Yamasaki *et al.* 1999].

The biological effects attributed to NO primarily arise because of the molecules ability to oxidise the sulphydryl groups of proteins to form S-nitrothiols via S-nitrosylation of cysteine residues [Simon *et al.* 1996; So *et al.* 1998]:-

$$2 \text{ NO} + \text{R-S-S-R} \rightarrow 2 \text{ R-S-NO}$$
(3)

For example, S-nitrosylation of c-Jun N-terminal kinase 2 [So *et al.* 1998] and upto seven different caspases [Jianrong *et al.* 1997] can reversibly inhibit these enzymes thereby rescuing animal cells that were undergoing programmed cell death.

The reaction between the superoxide anion  $(O_2)$  and NO to form the highly reactive and therefore very toxic peroxynitrite radical (OONO) is only limited by diffusion [Beckman *et al.*1994] and only prevented by the anti-AOS enzyme superoxide dismutase:-

$$NO + O_2 \rightarrow OONO$$
 (4)

The peroxynitrite radical quickly establishes an acid-base pair in mildly alkaline and neutral solutions thus forming peroxynitrous acid (OONOH) and can rapidly react with any dissolved carbon dioxide (CO<sub>2</sub>) to form the nitrosoperoxocarbonate adduct (OONOCO<sub>2</sub>):-

$$\begin{array}{ccccccccc} OONO^- & + & H^+ & \rightarrow & OONOH & (5) \\ OONO^- & + & CO_2 & \rightarrow & OONOCO_2^- & (6) \end{array}$$

The formation of the peroxynitrite and nitrosoperoxocarbonate radicals also leads to the oxidation of protein bound sulphydryl groups in a similar, but more reactive manner to NO [Beckman *et al.* 1994; Radi *et al.* 1991], however, these radicals can additionally inactivate proteins via their reactions with the catalytic iron-sulphur centres of metalloproteins [Stamler *et al.* 1992; Castro *et al.* 1994] to form iron-nitrosyl complexes that are generally inhibitory, eg: plant cell wall peroxidases [Ferrer & Ros Barcelo 1999].

The strong oxidant properties of the peroxynitrite radical are well documented [Gatti *et al.* 1994; van der Vliet *et al.* 1994; Ischiropoulos *et al.* 1995] and in fact the oxidation/nitration of tyrosine residues within proteins is considered to be a specific marker of peroxynitrite presence [Gow *et al.* 1996] in the same manner that protein-carbonyl formation is indicative of AOS, and specifically the hydroxyl radical:-



This specific protein modification induced by the peroxynitrite radical can prevent the deliberate enzymatic phosphorylation of tyrosine residues and consequently has the potential to modulate protein activity, for example, Kong *et al.* [1995] have shown that peroxynitrite mediated nitration of a single tyrosine residue of the cell cycle kinase, cdc2 ultimately blocks tyrosine phosphorylation and thus alters the relative duration of the cell cycle phases. This is a novel and potentially substantial protein controlling mechanism because upto 130 individual proteins can be nitrated at tyrosine residues by chemically generated peroxynitrite [Nielsen 1995]; whether the same process is occurring *in vivo* is unknown. The nitration of organic ring structures is not purely limited to protein bound tyrosine residues, and in fact, many plant phenolic compounds have been experimental used as antioxidants against the action of the peroxynitrite radical [Sandova *et al.* 1997].

The exact roles that NO plays in plants have not yet been clearly identified because of the relative infancy of the field, but some generalisations and inferences can be made between the more advanced animal studies and the scant plant data.

In animal systems intracellular NO signals are typically designated as cyclic guanylate monophosphate (cGMP) dependent, because a major effect of NO in animal cells is the activation of the soluble, cytosolic guanylate cyclase enzymes via iron-nitrosyl formation at the active site or S-nitrosylation of cysteine residues [McDonald & Moss 1994]. cGMP has been unequivocally discovered in plant cells [Brown & Newton 1992] and NO treatment has been shown to affect the cytosolic levels of cGMP [Pfeiffer *et al.* 1994] furthermore, cGMP has been proven to induce the transcription of genes, including some pathogenesis-related proteins and phenylalanine ammonia lyase [Durner *et al.* 1998] and to alter the production of secondary metabolites [Bowler *et al.* 1994]. All of these data

point towards similar roles of NO in plants as have been discovered in animals, but it should be noted that no guanylate cyclase enzymes or genes have yet been identified in plants, therefore the data should be interpreted tentatively.

NO can readily traverse phospholipid membranes and has therefore been touted as a potential extracellular molecule. Durner *et al.* [1998] have even proposed that NO may act as the long distance messenger molecule that triggers the acquisition of systemic resistance because of similarities with animal systems. Systemic acquired resistance requires the pathogen induced movement of a phloem-mobile signal to move from the point of infection systemically through the plant [Dempsey *et al.* 1999]. In mammals NO circulates in the blood as S-nitrothiol adducts of serum albumin and haemoglobins or as low molecular weight nitrosoglutathione (GSNO) the latter is believed to act as an effective intra- and inter-cellular NO carrier (see figure 1.5) [Jia *et al.* 1996]. What is intriguing is that glutathione is present in the apoplast [Polle *et al.* 1990] and is also a major constituent of the phloem, furthermore, GSNO has been found to be an extremely powerful inducer of plant defence responses [Durner *et al.* 1998].

In animal systems, AOS act synergistically with NO to induce programmed cell death [Brüne *et al.* 1998] and to directly execute microbial pathogens inside the macrophage vacuole [Mayer & Hemmens 1997], via the formation of the peroxynitrite radical. A similar programmed cell death response is known to occur during the pathogen-induced hypersensitive response in plants, and a parallel can once again be drawn to animal systems because this response is compromised by nitric oxide synthase inhibitors and mimicked by the addition of endogenously added peroxynitrite [Delledonne *et al.* 1998; Durner *et al.* 1998]. These results together with the recent discovery of homologues of the mammalian NADPH-oxidase complex in plants [Scheel 1998] reveal that very similar systems in both plants and animals that are responding to pathogen induced stress by the production of AOS and NO.

## **SECTION 2**

## **PROJECT OBJECTIVES & TIMELINE**

## 2.1 Project Objectives

The ultimate objective of this project was to utilise relatively simple techniques such as vacuum infiltration to study the leaf apoplast in plants. The overall proposed outcome was the identification of <u>changes</u> in the leaf apoplast. This project did not set out to explore any areas of apoplastic carbohydrate biochemistry, and was specifically focused on the quantitative and qualitative assessment of changes in the protein complement and enzyme activity of the apoplast and the generation of apoplastic radicals in the leaf apoplast in response to developmental changes and abiotic stress.

The initial position was therefore to provide a suitable apoplastic 'standard' to measure change, and from this, it was established that it would be necessary to optimise the original small-scale vacuum infiltration strategy adopted by previous students. Furthermore, because this strategy yielded low quantities of apoplastic extract it would be pertinent to develop a larger-scale, bulk procedure to selectively extract apoplastic components. It was hoped that the following specific questions would be answered during the course of the work: -

- 1. An initial identification and cataloguing of tobacco leaf apoplastic proteins using 1D electrophoresis.
- 2. The establishment of a complementary analysis of apoplastic proteins using a 2D system of electrophoretic separations to produce, 'protein maps'.
- 3. Using the above systems to identify proteins that change in response to stress and development.
- 4. The identification and cataloguing of enzymes present in the apoplast.
- 5. The identification of radicals either generated by, or released into the apoplast during stress.

It was hoped that N-terminal protein sequencing and database searches would be utilised to identify specific protein components. Identification of individual protein components involved during developmental processes and in response to stress would either fall within current models of the roles of the apoplastic compartment or shed light on 'new' apoplastic processes.

From conception, the need for highly resolving gel techniques had been identified as crucial to resolving small changes in apoplastic protein components. 2-dimensional (2D) electrophoresis is the most highly resolving of all electrophoretic techniques. Consequently, a major early objective of this project was to achieve good 2D separation of apoplastic protein components hopefully for further identifying analysis. Any 2Dsystem developed had to reflect the requirements of the project, in that the system had to cope with a high enough protein loading so that N-terminal sequencing could be attempted from individual 2D gels. N-terminal sequencing was chosen as the preferred method for protein identification because of the huge availability of database sequences for comparison. MALDI-TOF mass spectrometry was considered but the lack of comparable databases for sequence searches and access to suitable equipment dissuaded further development of this approach. Attempting to develop a 2D system with high protein loading is a relatively difficult challenge as the first dimension of 2D is typically very sensitive to protein load. Consequently, it was envisaged early in the project that the development of such a 2D-system would take up a large part of the project.

During the course of the project it was identified that many stresses whether caused by biotic or abiotic processes, ultimately result in the production active oxygen species (AOS), and when concentrations of such species are high enough result in the phenomenon of oxidative stress. Superoxide dismutase, peroxidase and catalase have been identified in many reports as the rudimentary and fundamental enzymes involved in resistance and tolerance to oxidative stress. The apoplast has been identified as a compartment were AOS are liberated in response to certain developmental stimuli and in response to stress. Consequently, analysis of apoplastic materials for these enzymes and radicals was a major focus of this project.

Some of the work within this thesis provides an excellent starting point for further work on apoplastic processes.

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The diagram below indicates the approximate time allocated to each major area of study in this project.

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YEAR 1

## **SECTION 3**

## MATERIALS

## 3.1 Glassware and plasticware

All glassware used for protein manipulations was washed in a dishwasher using commercial detergents and purified water rinses. Prior to use glassware was rinsed with purified water (Elgastat, UHQPS, Elga Ltd, Buckinghamshire, UK) and allowed to dry. Microcentrifuge tubes, pipette tips, cuvettes and 96 well microtitre plates were obtained from Sarstedt Ltd (Leicester, UK). Petri dishes were supplied by Bibby Sterilin Ltd (Staffordshire, UK).

## 3.2 Plant material

*Nicotiana tabacum* (SR-1) seeds were supplied by Dr R.R.D. Croy (Department of Biological Sciences, University of Durham, UK). Potato varieties were supplied by the Potato Research Council, Sutton Bridge, Lincs., UK.

## 3.3 Buffers and solutions

All buffers and solutions were prepared with purified (Elgastat, UHQPS, Elga Ltd, Buckinghamshire, UK) or distilled water.

## 3.4 Chemical and biological reagents

Chemical reagents were supplied by BDH Ltd (Leicestershire, UK) and were of AnalaR grade. Other reagents were obtained from the following sources: -

Sigma (Poole, Dorset, UK)

Acrylamide, ammonium persulphate, bis-acrylamide, Bovine serum albumin, bromophenol blue, Coomassie brilliant blue R250 CI 42660, dithiothreitol, DNAse, glycerol, molecular weight markers (MW-SDS-70L and MW-SDS-200), RNAse A, SOD from bovine erythrocytes, sodium dodecyl sulphate, TEMED, Tween-20, XTT, NBT, PMS.

 BioRad (Hemel Hempstead, Hertfordshire, UK) Bradford protein assay kit, silver stain kit. • National Diagnostics (Hull, UK)

Protogel, 30%(w/v) acrylamide: 2.7%(w/v) cross-linked with methylene bisacrylamide (37.5:1 ratio) in distilled water, ammonium persulphate.

- Amersham Pharmacia, Uppsala, Sweden
   Carrier ampholytes (ranges 5-7 & 3-10); native IEF marker proteins; carbamylation standards.
- Micron Separations (MA, USA)
   Problott PVDF transfer membrane.

## **SECTION 4**

## METHODS

## 4.1 Plant growth and potato storage conditions

Seeds of SR-1 tobacco (*Nicotiana tabacum*) were sown in trays containing Levington M3 compost (Levington, UK). The seedlings that had emerged after 14 days were transferred to 2 litre pots and grown in a controlled environment room at 23/20  $\pm$  2°C day/night temperatures, 16 hour photoperiod (20:00-12:00 h), 100µE s<sup>-1</sup> m<sup>2</sup> (50 W m<sup>2</sup>) light intensity (cool-white fluorescent tubes) and 70  $\pm$  5% relative humidity. The plants were fertilised weekly with 1% (w/v) Phostrogen (Phostrogen, UK) and watered once every three days.

Potatoes of various varieties were either bought from a local supermarket or supplied by the potato research council from Sutton Bridge, Lincolnshire. Potatoes were routinely stored in lightproof containers at 10°C. For the 24-hour period before potatoes were exposed to impact stress they were moved to lightproof containers at 4°C.

## 4.2 Extraction and preparation of apoplastic components

The principle of vacuum infiltration (VI) to recover apoplastic components was essentially the same as that described by Hammond-Kosack [1992] with adaptations made by Dr C. llett, Dr R. Croy and during the course of this study. Initially air present in apoplastic spaces was removed by the application of a vacuum to the surface of a liquid into which the tissue had been submerged. The slow release of the vacuum caused the liquid to flood into the apoplast (see the working definition of the term apoplast in section 1.3.4) after which the intercellular fluid (ICF) was collected by low speed centrifugation. The ICF collected contains polysaccharides, apoplastic proteins and a range of other components [Hammond-Kosack 1992]. A small scale technique had been in use by a previous student but it was only capable of isolating small volumes of ICF. Therefore, a large scale technique was developed to extract larger quantities of ICF, the developmental process is outlined in section 5. Both techniques were used in tandem to extract ICF during the project, these methods are described in sections 4.2.1 and 4.2.2. To ensure that the ICF recovered is uncontaminated by intracellular components, comparisons had to be made between it and the total soluble proteins (TE's) present in the leaf, the extraction process for TE's is detailed in section 4.2.3. ICF is typically very dilute, consequently, several approaches were taken to concentrate ICF for further analysis, these are described in sections 4.2.4 and 4.2.5.

## 4.2.1 Small scale VI protocol

Upto 50 tissue strips of approximately 2cm<sup>2</sup> were cut from leaves with a sharp blade. Care was taken to avoid the major veins, tissue strips were then washed for >15 minutes in  $4^{\circ}$ C-chilled dH<sub>2</sub>O. The tissue was then dried off using paper towel and then submerged under 4°C-chilled infiltration buffer (200mM NaCl, pH 7.5) in a vacuum jar using a wire mesh and 1 kg weight. The gas was removed from apoplastic spaces by the application of a vacuum (8.5 x  $10^4$  Pa (Nm<sup>-2</sup>)) for 75 seconds using a KNF Neuberger pump (model number NO35AN) attached to the vacuum jar, see figure 4.1 [A]. The vacuum jar was gently agitated throughout gas removal so that bubbles were dislodged from the tissue surfaces. The apoplast was then flooded by slow release of the vacuum over a 15-second period. Each leaf strip was then individually dried on paper towel to remove any excess buffer, carefully rolled and inserted into a 0.5-ml eppendorf tube with a small hole cut in the bottom; this arrangement was then placed inside an intact 1.5-ml eppendorf and centrifuged (Beckmann GS-15R, F3602 rotor, 4°C, 900g) for 10 minutes. Any ICF that had a green colouration was discarded, (because it indicated cellular damage and therefore intracellular leakage), the remaining ICF was pooled and either stored at -20°C or dialysed overnight against dH<sub>2</sub>O at 4°C, lyophilised and stored at -20°C in a parafilm sealed container (see 4.2.4). A small aliquot (<100µl) of ICF was retained after every VI and assayed for malate dehydrogenase-activity, this was used to calculate the degree of intracellular contamination of ICF (see section 4.3.1) and if required a Bradford assay was also undertaken to quantify the protein concentration (see section 4.3.2). The Bradford assay was only carried out on small scale recovered ICF because large scale ICF were too dilute.

## 4.2.2 Large scale VI protocol

Leaf tissue strips were prepared, washed and dried as described above (section 4.2.1) - the major difference here is the quantity of tissue, the large scale method permits VI to be carried out on upto 200 tissue strips at a time. The tissue strips were then arranged along a length of nylon mesh (10cm x 1m – supplied by Lockertex, UK) and rolled to form a 'Swiss roll' taking care not to crush or tear the tissue. The Swiss-roll was held in place with a loosely fitting elastic band and infiltrated as described as above (section 4.2.1), excess buffer was allowed to drain from the tissue and nylon mesh after wrapping each roll in tissue paper and standing end up. Each Swiss-roll was then inserted into a sterile blood sample tube with a 3-4 mm hole cut in the bottom; the blood sample tube was then placed into an intact 50-ml Falcon tube. Figure 4.1 [B]-[D] illustrates the large scale VI method. The assembly was then subjected to centrifugation at 2000 rpm for 10 minutes (4°C, Sigma-204 centrifuge, 50ml swing-out rotor) permitted recovery of ICF. Any ICF that had a green colouration was discarded, (because it indicated cellular damage and



# Figure 4.1

Photographs of the equipment and methodology of the large scale VI method

[A] vacuum pump and jar utilised for VI protocols described in sections 3.6.1 and 3.6.2; [B] tobacco leaf strips arranged onto nylon mesh; [C] rolled nylon mesh-tissue strip 'Swiss' roll inside sample tube; [D] the whole assembly is inserted into a 50-ml Falcon tube. therefore intracellular leakage), the remaining ICF's were pooled and either stored at –  $20^{\circ}$ C or dialysed overnight against dH<sub>2</sub>O at 4°C, lyophilised and stored at - $20^{\circ}$ C in a parafilm sealed container (see 4.2.4). A small aliquot (<1ml) of ICF was retained after every VI and assayed malate dehydrogenase-activity, this was used to calculate the degree of intracellular contamination of ICF (see section 4.3.1).

## 4.2.3 Total soluble protein extraction

For comparisons between apoplastic and intracellular proteins to be made total soluble protein extracts (TE's) were required. TE's also formed the basis for calculating relative malate dehydrogenase activity (see section 4.3.1). All leaves selected for TE preparation were healthy, clean, dry and developmentally and physiologically identical (as judged by the investigator) to those used in the ICF preparation. 2cm<sup>2</sup> strips of leaf tissue were excised, taking care to avoid major leaf veins and then weighed. The tissue was then around to a fine powder in a pestle with quartz sand and liquid  $N_2$ . When the  $N_2$  had boiled off buffer was then added to the ground tissue (the same as utilised for VI plus 1 mM DTT) at a ratio of 1.5-ml per 1 gram of original tissue, this mixture was then ground. The resultant slurry was aliquoted into 1.5 ml eppendorf tubes and centrifuged (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes at 4°C to pellet out sand and large cellular debris. The resultant supernatants were then decanted, pooled and ultracentrifuged at 50,000 rpm, 4°C for 60 minutes. The final supernatant was aliquoted into eppendorf tubes (1ml volumes), the protein concentration estimated (section 4.3.2) and the malate dehydrogenase activity assayed (section 4.3.1). Aliquots were stored at -20°C. If thought necessary for comparative purposes, TE was dialysed overnight against 4°C-chilled dH<sub>2</sub>O, lyophilised and the solids stored in parafilm sealed containers at -20°C (section 4.2.4).

## 4.2.4 Lyophilisation of recovered ICF and TE

Typically ICF has a very low protein concentration, dialysis, to remove salts from the apoplast and those used in the infiltration buffer, followed by lyophilisation was a useful method to concentrate apoplastic components and was used throughout this project. Dialysis was carried out immediately after ICF had been recovered by centrifugation (sections 4.2.1 & 4.2.2), using narrow width dialysis tubing ( $M_r$  cut-off 10 kDa from Sigma, UK) against 2-3 changes of dH<sub>2</sub>O for a minimum of 8 hours, at 4°C. The material was recovered after dialysis and assayed for protein content using the Bradford assay (section 4.3.2). According to the Bradford assay results appropriate aliquots (200-800µl containing 100µg of protein material) of dialysed ICF were placed into 1.5ml-eppendorf tubes with 2-3 small holes piercing the lid. Samples were quickly frozen in liquid N<sub>2</sub> and

placed into the freeze-drier overnight and lyophilised. Dried solids were stored in parafilm sealed eppendorf tubes at -20°C until required.

## 4.2.5 Acetone precipitation of recovered ICF and TE

Acetone precipitation was utilised early in the project to concentrate ICF, but was later abandoned because of the potential of acetone to N-terminally block proteins, this would ultimately reduce the number of proteins that could effectively be sequenced, 100% acetone,  $-20^{\circ}$ C cooled was added to ICF at a ratio of 2:1 and mixed; proteins were allowed to precipitate for 15 minutes at  $-20^{\circ}$ C. Precipitates were then separated out by centrifugation (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes. The supernatant was discarded and the pellet of precipitated proteins was washed in  $-20^{\circ}$ C chilled 70% (v/v) acetone at a ratio of 2:1 of the original volume of sample. Following a repeat 5-minute centrifugation the supernatant was discarded and the pellet was air-dried for 20 minutes. Dried solids were stored at  $-20^{\circ}$ C until required.

## 4.3 Assay analysis of ICF and TE

To assess the degree of intracellular contamination caused by cellular damage and leakage, all ICF's & TE's were routinely assayed for the activity of the cytoplasmic/mitochondrial enzyme malate dehydrogenase (section 4.3.1). When required the Bradford assay (section 4.3.2) was used to provide an estimate of protein concentration in all samples. Other assays were performed in order to assess the extracellular activities of enzymes of interest (see sections 4.3.3 to 4.3.6).

## 4.3.1 Malate dehydrogenase (MDH) assay for estimating intracellular contamination of ICF preparations

Malate dehydrogenase (MDH, EC 1.1.1.37) activity was assayed in both TE's and ICF's by a method described in Biochemica Catalogue Enzymes [1968]. A reaction cuvette containing 943 $\mu$ l of 0.1 M phosphate buffer, pH 7.5, 33 $\mu$ l of 2mg/ml oxaloacetate in phosphate buffer, 17 $\mu$ l of 10 mg/ml NADH (sodium salt) in dH<sub>2</sub>0, was first zeroed against a duplicate mixture. The assay was initiated by the addition of 7 $\mu$ l of either TE or ICF to the reaction cuvette which was quickly inverted twice to mix the reagents and sample. The reaction was followed by measuring the decrease in absorbance at 340nm for 60 seconds. In order to 'quantify' the degree of intracellular contamination of the ICF, TE readings were normalised to 100% of the MDH-activity present in the tissue [Ilett 1998]: -

 $\Delta Ab_{340}$  of ICF x100 = Relative MDH-activity  $\Delta Ab_{340}$  of TE Only ICF's with <1% relative MDH-activity were used for apoplastic analysis.

## 4.3.2 Bradford Assay for the quantification of protein

The spectrophotometric dye-binding assay based upon the work of Bradford [1976] and Spector [1978] was used to estimate protein concentration in all samples. The principle of the assay is based on the observation that the maximal absorbance for an acidic solution of Coomassie brilliant blue G-250 shifts from 465nm to 595nm when protein binding occurs. Bradfords' reagent was purchased from BioRad (Poole, Dorset, UK).

In order to quantify the total protein content of an unknown sample a series of standard bovine serum albumin (BSA) solutions ranging from 0.5µg to 10µg per assay volume were used to produce standard curves. 100µl volumes of sample, standards and water blanks were added to 900µl of Bradfords' reagent in micro-cuvettes and allowed to stand at room temperature for 7 minutes. Absorbencies at 595nm were measured using the Unicam UV2 spectrophotometer (ATI Unicam, Cambridge, UK) after zeroing using the water blanks, standard curves and quantification of unknown samples was accomplished using Prism GraphPad software (Prism Software, USA).

For speed of throughput a microtitre plate version of the above assay was established. A 1mg/ml stock solution of BSA in infiltration buffer (see sections 4.2.1 and 4.2.2) was used to produce a range of calibration standards between 100 and 1000µg/ml. ICF and TE samples are diluted as necessary with infiltration buffer. 10µl of each standard and an appropriate volume of ICF or TE sample was added and each well was then diluted by the addition of infiltration buffer to a total volume of 160µl. 40µl of Bradford reagent was added to each well and the plate is shaken for 30 seconds and left to incubate for a further 7 minutes. After the incubation the absorbance of each well was measured against an infiltration buffer blank using the Dynatech MR5000 microtitre plate reader (Dynatech, UK) at 570nm. The quantification of unknown samples and the construction of the calibration curve was accomplished using Prism GraphPad software (Prism Software, USA).

## 4.3.3 Superoxide Dismutase Assay

Superoxide dismutase (SOD) activity in ICF's was quantified using an assay developed during the course of this research [see memorandum, page iii]. A series of SOD calibration standards were prepared from a commercial preparation of SOD (Sigma, Poole, Dorset, UK), dissolved in 50mM phosphate buffer, pH 8.2. The SOD calibration standards (ranging from 0.01U (2.875ng) to 100U (28.75µg)), experimental samples (prepared in 50mM phosphate buffer, pH 8.2) and blanks (50µl of 50mM phosphate

buffer, pH 8.2) were added to the wells of a microtitre plate in a 25µl aliquot, in triplicate. 200µl of 50mM phosphate buffer, pH 8.2, containing 0.1mM EDTA, 98µM NADH, 150µM 2,3-bis-[2-methoxy-4-nitro-5-sulphophenyl]-2H-tetrazolium-5-carboxanilide (XTT) was then added to each well. The reaction was initiated with 25µl of a 28µM solution of phenazine methosulphate (PMS) in 50mM phosphate buffer, pH 8.2. This mixture had final concentrations of 80µM EDTA, 78.4µM NADH, 120µM XTT and 2.8µM PMS. Absorbance readings were taken at  $\lambda_{max}$  (450nm) for 2 minutes from which the mean blank absorbance value was initially subtracted and then the adjusted initial rate ( $\Delta_{450}$ /min) was calculated using data derived from the Dynatech MR5000 microtitre plate reader (Dynatech, UK). A calibration curve of known SOD units (calibration standards) versus  $\Delta_{450}$ /min was constructed and quantification of experimental samples was achieved by reading initial rate values from this standard curve (Prism GraphPad software - Prism Software, USA).

## 4.3.4 Guaiacol Peroxidase Assay

The activities of peroxidases (POD's) that can utilise the artificial substrate, guaiacol were quantified in ICF. The reaction mixture comprised of 2.9ml of 0.1 M phosphate buffer, pH 7.0, 50 $\mu$ l of 0.0022% (v/v) guaiacol in 0.1 M phosphate buffer, pH 7.0, and 40 $\mu$ l of 0.03% (v/v) hydrogen peroxide in 0.1 M phosphate buffer, pH 7.0. The reaction mixture was initially zeroed against a duplicate mixture and the assay begun by adding 10 $\mu$ l of an ICF preparation (prepared in 0.1 M phosphate buffer, pH 7.0) to the reaction cuvette which was then quickly inverted twice to mix the reagents and sample. The reaction was followed by measuring the increase in absorbance at 470nm using a Unicam UV2 spectrophotometer (ATI Unicam, Cambridge, UK). Experimental sample enzyme activities were then calculated either from the published extinction coefficient for guaiacol of 26.6 x 10<sup>6</sup> M<sup>-1</sup> cm<sup>-1</sup> [George 1953] or from a calibration curve of known POD standards and reading unknowns from the curve (Prism Software, USA).

## 4.3.5 Superoxide Anion quantification from tissue

Tuber tissue was excised from potatoes as small cores (~5 mm diameter x 3 mm long) weighing in the order of 30-35mg each. After extensive washing with dH<sub>2</sub>O to remove any intracellular materials that would have been released during core excision, the tuber cores were each placed into a 200 $\mu$ l volume of 50mM phosphate buffer, pH 8.2 containing, 120 $\mu$ M XTT in an eppendorf tube. The mixture was incubated for 20 minutes in the dark, at room temperature after which the tuber core tissue was removed and the solution was centrifuged (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes to pellet any cell debris that may interfere with subsequent absorbance measurements. The supernatant was withdrawn and the absorbance measured at 450nm against a reagent-

tissue blank using the Beckman DU7500 (Beckman Instruments, UK). Superoxide anion concentration was calculated using the published extinction coefficient for XTT, of 2.16 x  $10^4$  M<sup>-1</sup> cm<sup>-1</sup> [Sutherland & Learmonth 1997].

## 4.3.6 Nitric Oxide quantification from tissue

Tuber cores were prepared as described in section 4.3.5 and after extensive washing with dH<sub>2</sub>O, the tuber cores were each placed into a 200 $\mu$ l volume of 50mM phosphate buffer, pH 7.8, containing 114mM sulphanilamide (SULF) and 2.4mM N-(1-napthyl)-ethylenediamine dihydrochloride (NEDD). The mixture was incubated for 20 minutes in the dark, at room temperature after which the tuber core tissue was removed and the solution was centrifuged (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes to pellet any cell debris that may interfere with subsequent absorbance measurements. The supernatant was withdrawn and the absorbance measured at 496nm against a reagent-tissue blank using the Beckman DU7500 (Beckman Instruments, UK). Nitric oxide concentration was calculated using the published extinction coefficient for the SULF-NEDD reagent, of 1.25 x 10<sup>4</sup> M cm [Nims *et al.* 1995].

## 4.4 1D gel analysis strategies

ICF and TE preparations were subjected to 1D gel analysis for visualisation of protein banding patterns (sections 4.4.1 to 4.4.4), in gel enzyme activity assessments (sections 4.4.5 to 4.4.7), for subsequent N-terminal sequencing (section 4.4.11), antibody probing (section 4.4.9) or oxyblotting for the detection of oxidatively modified protein components (section 4.4.10).

## 4.4.1 SDS-PAGE

Polypeptides were size fractionated by SDS-PAGE performed essentially as described by Laemmli (1970). 1.5 to 0.75 mm-thick vertical slab gels were cast and electrophoresed using the BioRad Mini-Protean II system. Unless otherwise stated, 12% (w/v) acrylamide/bis-acrylamide (37.5:1) separating gels (containing 375mM Tris-HCI, 0.1% (w/v) SDS) and 4% (w/v) acrylamide/bis-acrylamide stacking gels (containing 125mM Tris-HCI, 0.1 (w/v) SDS) were used throughout these studies. Liquid sample (i.e.: TE's) were prepared by diluting the sample with 2x sample loading buffer (125 mM Tris-HCI pH6.8, 6%(w/v) SDS, 0.01% (v/v)  $\beta$ -mercaptoethanol, 10% (v/v) glycerol, 0.05% (w/v) bromophenol blue) at a ratio of 1:2. Solid protein material (i.e.: ICF's) was reconstituted in a suitable volume of 1x sample loading buffer. Samples were denatured just prior to loading onto a gel by incubating for 5 minutes in a boiling heating block. Electrophoresis was carried out with 1x running buffer (3 g/l Tris-HCI pH 8.3, 14.4 g/l glycine, 1 g/l SDS), initially at 70V until the marker dye front had just entered the separating gel, then at 170V until the dye front had reached the bottom of the gel.

## 4.4.2 Native-PAGE

Proteins were separated based on three physical properties, electrical charge, overall mass, and shape by native-PAGE. The native-PAGE protocol used was essentially an adapted non-denaturing version of the SDS-PAGE protocol presented in section 4.4.1 0.75 mm-thick vertical slab gels were cast and electrophoresed using the BioRad Mini-Protean II system. Unless otherwise stated, 12% (w/v) acrylamide/bis-acrylamide (37.5:1) separating gels (containing 375 mM Tris-HCl) and 4% (w/v) acrylamide/bis-acrylamide stacking gels (containing 125 mM Tris-HCl) were used throughout these studies. Liquid sample (i.e.: TE's) were prepared by diluting the sample with 2x native sample loading buffer (125 mM Tris-HCl pH6.8, 10% (v/v) glycerol, 0.05% (w/v) bromophenol blue) at a ratio of 1:2. Solid protein material (i.e.: ICF's) was reconstituted in a suitable volume of 1x native sample loading buffer. Samples were not heated. Electrophoresis was carried out with 1x running buffer (3 g/l Tris-HCl pH 8.3, 14.4 g/l glycine), initially at 70V until the marker dye front had just entered the separating gel, then at 170V until the dye front had reached the bottom of the gel.

## 4.4.3 Coomassie based protein staining

Protein bands were visualised by submerging gels in 0.1% (w/v) Coomassie blue R250, in 40% (v/v) methanol, 10% (v/v) glacial acetic acid, and destained with 40% (v/v) methanol, 10% (v/v) glacial acetic acid. Alternatively a more sensitive Coomassie based stain was also utilised, 'Fast Stain' from Zoion, (Mass., USA), following the manufacturers instructions.

## 4.4.4 Silver staining of proteins

Protein bands were visualised by silver staining with the BioRad silver staining kit following manufacturers instructions. Briefly, the rapid silver staining protocol was adopted following the sequence of; submerging the gel in fixative (40% (v/v) methanol, 10% (v/v) glacial acetic acid) for >30 minutes; submerging the gel in oxidiser for 5 minutes; washing the gel extensively with  $QH_2O$  for 15 minutes; submerging the gel in silver reagent for 20 minutes; washing the gel with  $QH_2O$  for 1minute; developing the bands by submerging in developer reagent; and finally, stopping development by submerging the gel in 5% (v/v) acetic acid solution.

## 4.4.5 In gel activity staining for catalase

After native electrophoresis (section 4.4.2) the gel was removed from the cassette, submerged into a substrate solution comprising of either 3% (v/v) or 0.3% (v/v) hydrogen peroxide, and incubated in the dark with agitation for 15 minutes. The solution was then removed and the gel washed for >1 minute with dH<sub>2</sub>O. The gel was then submerged in the staining solution which comprised 1% (w/v) potassium ferricyanide and 1% (w/v) ferric chloride. Yellow to pale blue bands of catalase activity should appear on a blue background.

## 4.4.6 In gel activity staining for peroxidase

After native electrophoresis (section 4.4.2) the gel was removed from the cassette and submerged into a staining solution of 0.0225% (v/v) hydrogen peroxide and 0.001% (v/v) guaiacol in 50mM sodium acetate buffer at pH 5.0. The gel was incubated in the dark until peroxidase bands of activity were stained red-brown.

## 4.4.7 In gel activity staining for superoxide dismutase

After native electrophoresis (section 4.4.2) the gel was removed from the cassette and submerged into a staining solution of  $0.98\mu$ M nitroblue tetrazolium,  $1.57\mu$ M phenazinemethosulphate and  $5.9\mu$ M magnesium chloride contained in 50mM Tris-HCl at pH 8.5. The gel was incubated under direct light (a lamp is ideal) while being agitated. Superoxide dismutase activity bands were detected as clear zones on a dark blue background.

## 4.4.8 Gel drying for storage

After any of the treatments detailed above gels were routinely dried down for storage. The procedure was based on the protocol provided with the BioRad gel drying apparatus and consisted of, incubating the gel for >1 hour in many changes of  $dH_2O$ . Rehydrating sheets of gel wrap in water and sandwiching the gel between 2 sheets of gel wrap using the gel-drying frame as described in the manufactures instructions. The gel-drying frame was placed into the drying apparatus until gels were dehydrated.

## 4.4.9 Electroblotting from gels for antibody probing

After electrophoresis was complete the enclosing glass plates were removed and the gel was submerged in 1x transfer buffer (0.965g/l Tris base, 4.5g/l glycine) for 15-20 minutes. Suitably sized pieces of nitrocellulose membrane (Micron Separations, MA,
USA) were cut and wetted in methanol for 2 minutes and then placed into 1x transfer buffer for 5 minutes to equilibrate. Four pieces of 3MM blotting paper were cut to the same size as the nitrocellulose membrane and dampened with 1x transfer buffer. The BioRad Mini-Protean II Wet Blotting system was utilised to effect protein transfer, the manufacturers instructions were followed. Briefly, the two electroblotter sponges were submerged in 1x transfer buffer and squeezed to exclude as much trapped air as possible. The electroblotter unit was assembled as described in the manufacturers instructions and inserted into the tank. Electroblotting was undertaken with 1x transfer buffer for 30-60 minutes at 100 V. The unit was then disassembled, the supporting matrix removed, and stained briefly with Coomassie or Ponceau S to ensure protein transfer. It was usual to stain the gel to ensure that protein transfer had been efficient.

## 4.4.10 Oxyblot analysis

The oxyblotting kit was purchased from Oncor (Oncor, France) and enabled the detection of oxidatively modified proteins, specifically those modified by the action of the hydroxyl radical. The kit was used essentially as described by the manufacturers.

Briefly, the protein solution to be oxyblotted was derivatised with an acidic solution of dinitrophenylhydrazine (DNPH), this procedure induces DNPH-protein complexes at sites of hydroxyl-radical mediated oxidation, specifically carbonyl groups. The derivatisation reaction was stopped after 5 minutes by the additon of 2M Tris-base (neutralisation solution). The protein solution was then loaded onto a standard SDS-PAGE gel (see section 4.4.1) and electrophoresed. After electrophoresis the separated proteins were electroblotted from the gel (see section 4.4.9) and antibody probed (next paragraph).

Primary antibodies were provided by the manufacturer. The antibody probing procedure followed these steps:-

- 1. 45 minute incubation in blocking buffer (tris buffered saline (TBS) containing 3% non-fat dried milk).
- 2. 45 minutes in blocking buffer containing 1:150 anti-DNPH primary antibody.
- 3. 3 x 5 minute washes in TBS-Tween.
- 4. 45 minutes in blocking buffer containing alkaline phosphatase linked secondary antibody.
- 5. 3 x 5 minute washes in TBS-Tween.
- 6. Bands were visualised using Sigma BCIP/NBT tablets prepared as per manufacturers instructions.

## 4.4.11 Electroblotting from gels for N-terminal sequencing

Proteins were separated as in standard SDS-PAGE (see section 4.4.1) except that the gel was either cast at least 12 hours before use and stored, or pre-run for 30 minutes at 50V with  $200\mu$ M thioglycolic acid in the upper buffer chamber. These measures ensure that any reactive amines present in the gel matrix have either fully derivatised or are scavenged thereby lessening the chances of inadvertent N-terminal blocking of proteins.

After electrophoresis was complete the enclosing glass plates were removed and the gel was submerged in 1x transfer buffer (2.213g/l 3-(cyclohexylamino)-1-propanesulphonic acid, 10% (v/v) methanol, pH 11) for 15-20 minutes. Suitably sized pieces of PVDF (polyvinylidene fluoride) membrane (Micron Separations, MA, USA) were cut and wetted in methanol for 2 minutes and then placed into 1x transfer buffer for 5 minutes to equilibrate. Four pieces of 3MM blotting paper were cut to the same size as the supporting matrix and dampened with 1x transfer buffer. The BioRad Mini-Protean II Wet Blotting system was utilised to effect protein transfer, the manufacturers instructions were followed. Briefly, the two electroblotter sponges were submerged in 1x transfer buffer and squeezed to exclude as much trapped air as possible. The electroblotter unit was assembled as described in the manufacturers instructions and inserted into the tank. Electroblotting was undertaken with 1x transfer buffer for 30-60 minutes at 100 V. The unit was then dismantled, the supporting matrix removed, and stained briefly with Coomassie blue or Ponceau S to ensure efficient protein transfer had occurred.

## 4.5 2D gel analysis strategies

ICF and TE preparations were subjected to separation in a variety of different 2D-gel systems in an attempt to produce high resolution protein maps (see sections 4.5.6 and 4.5.7 for the most successful). Throughout these experiments ICF components continually precipitated and interfered dramatically with the separation therefore a variety of sample preparation techniques were employed to try and circumvent this problem. These are presented in sections 4.5.1 to 4.5.3. The isolation of ICF was extremely laborious and time-consuming and a more convenient control mixture of proteins isolated from mitochondria was available (section 4.5.4) therefore this preparation was used for the development of the 2D systems described in this thesis.

## 4.5.1 Standard TE and ICF sample preparation for 2D analysis

Known quantities of lyophilised or acetone precipitated TE (sections 4.2.4 and 4.2.5) or lyophilised ICF proteins (section 4.2.5) were reconstituted in a suitable quantity of isoelectric focussing (IEF) sample loading buffer (dependent upon IEF and 2D

technique). Precipitate was pelleted by centrifugation (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes prior to loading the sample onto the IEF.

## 4.5.2 ESA based TE and ICF sample preparation for 2D analysis

Samples were heat denatured by the addition of  $100\mu$ l of boiling 0.3% (w/v) SDS, 200mM DTT, 28mM Tris-HCl, 22mM Tris-base, followed by 5 minutes in a  $100^{\circ}$ C heating block. Samples were then cooled on ice for 5 minutes, treated with  $10\mu$ l of enzyme mixture (24mM Tris-Base, 476mM Tris-HCl, 50mM MgCl<sub>2</sub>, 1mg/ml DNAse I) and chilled for a further 10 minutes. Protein material was then removed from solution by incubating on ice for 20 minutes with ice cold acetone (80%(v/v)) followed by centrifugation at 15,000 rpm for 10 minutes, the resultant pellet was air-dried. Samples were then resuspended in a suitable volume of sample loading buffer (7.92M urea, 0.06% SDS, 1.76% ampholytes, 120mM DTT, 3.2% Triton X-100, 22.4mM Tris-HCl, 17.6mM Tris-Base).

## 4.5.3 Ammonium Sulphate-PVP TE and ICF sample preparation for 2D analysis

For TE preparation leaf tissue was initially cut up into small pieces, weighed and then placed into a 4°C-chilled blender. Liquid nitrogen was added and the tissue fragmented further with a clean spatula. At a ratio of 3ml per 1g of tissue 0.1 M phosphate buffer (pH7.5, 1mM DTT, 2mM EDTA) and at 1g per 10ml of buffer (i.e.: 10% (w/v)) insoluble PVP were then added. The mixture was blended for 2 minutes then strained through Miracloth into a beaker placed on ice and the volume was recorded. Ammonium sulphate was added spatula-wise with constant stirring to 80% relative saturation. Following centrifugation (4°C, 10,000 g, JA-14 rotor, Beckman J2-HC centrifuge) the supernatant was removed and spatula-sized quantities of pellet were collected and placed into eppendorf tubes. This ammonium sulphate stabilised preparation was routinely stored at -20°C for extended periods with no obvious detrimental effects. When required for analysis 1 spatula worth of ammonium sulphate stabilised TE was resuspended in 3-4ml of 0.1M Phosphate buffer (pH7.5 containing 1mM DTT and 2mM EDTA) and then microfuged for 2 minutes to remove any PVP. The supernatant was then decanted and a Bradford protein estimate obtained. A 2.5ml volume of sample was then loaded onto a QH<sub>2</sub>0 pre-equilibrated PD-10 column (Pharmacia Biotech) and once it had entered the Sephadex bed a further 3.5ml of QH<sub>2</sub>0 was added. The second and third millitre volumes off the column were collected and assayed to determine the protein concentration. 400µl centricons (spun at 4°C, 2000g, using the F2402 rotor in a Beckman GS-15R centrifuge for 60-120 minutes) were then utilised to reduce the volume and salt concentration and concurrently increase the protein concentration. Sample was then collected from the filter

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insert of the centricon, assayed for protein concentration, and was finally stored frozen at  $-20^{\circ}$ C until required for 2D-analysis.

ICF was collected as described in section 4.2.1 or 4.2.2, ammonium sulphate precipitation was not possible because ICF is very dilute. PVP was added at a concentration of 10% (w/v) and was removed after 30 minutes by centrifugation (benchtop, Eppendorf centrifuge at 15,000 rpm) for 5 minutes. The supernatant was collected, dialysed, lyophilised and then prepared as described in section 4.2.4.

## 4.5.4 2D standard control protein mixture (from blowfly mitochondria)

Control flies were grown at a constant  $24^{\circ}$ C. Flies were heat-shocked by incubating batches at  $37^{\circ}$ C for 40 minutes. After heat treatment, they were returned to a constant  $24^{\circ}$ C chamber for 4 hours to permit the heat-shock responses to develop. Blowfly muscle mitochondria were isolated from both control and heat shocked flies as described previously [Elwadawi *et al.* 1995]. Briefly, thoraces were collected and flight muscles removed and disrupted using a mortar and pestle. Mitochondria were centrifuged at 3,600 rpm (RCF = 1500g) in a Beckman Avanti 30 (F2402 rotor). The mitochondrial pellet was resuspended in 0.15M KCl, 1mM EDTA, 10mM Tris and stored frozen at –  $80^{\circ}$ C. Kind thanks are extended to Judith Chambers (Dept. of Biological Sciences, University of Durham) for the supply of this material.

## 4.5.5 Preparation of coloured pl markers

After many polyacrylamide-isoelectric focussing (PAG-IEF) trials that did not exhibit adequate protein separation a mixture of coloured proteins and a dye was concocted in order to quickly and visually assess the formation of the carrier ampholyte pH gradient. 6mg, 1mg and 3mg of commercial preparations of cytochrome C, myoglobin and the acidic dye methyl red were weighed on an analytical balance, respectively. Each was solubilised with the addition of 1ml, 800 $\mu$ l and 1ml of QH<sub>2</sub>0, respectively. The optimal mix of coloured pl markers was determined experimentally and the final combination used was 200 $\mu$ l of cytochrome C preparation, 100 $\mu$ l of myoglobin preparation, 20 $\mu$ l of methyl red preparation and 200 $\mu$ l of QH<sub>2</sub>0. A typical loading of this solution onto a PAG-IEF gel for visualisation was of the order 5-15 $\mu$ l.

## 4.5.6 In-house adaptation of ESA 2D system

This method is an in-house adaptation of the ESA/Oxford GlycoSystems Investigator<sup>™</sup> 2D electrophoresis system. 750µl aliquots of gel solution (9.5M urea, 2%(w/v) triton X-100, 4.1% acrylamide, 5mM CHAPS) were prepared and 46µl of 40(w/v) ampholytes and

 $5_{\rm ul}$  of 10% (w/v) APS were added and mixed thoroughly. Using a piece of rubber tubing attached to a 1ml syringe the tubing was secured to a suitably dimensioned capillary tube (1.7mm id x 17cm or x 8.5cm) and the free end placed into an eppendorf containing the gel solution. The gel solution was carefully draw up the capillary so that the gel was the desired length (6.5 or 15cm) polymerisation was then allowed to proceed for 60 minutes. Once polymerised the syringe and tubing were removed and any excess gel was cleaned from the bottom of the capillary tube. Anodic buffer (10 mM phosphoric acid) filled the lower chamber of the electrophoresis tank and the capillary gels were inserted into the apparatus ensuring that the bottom of the gels were in full contact with the buffer. The upper chamber was then filled with cathodic buffer (100 mM NaOH) and the ends of the gels were washed several times with buffer using a Hamilton syringe. Sample is prepared and loaded onto the cathodic end of the gels in a 10-30µl volume using a Hamilton syringe and 10<sup>ul</sup> of sample overlay solution (0.5M urea, 0.2%(w/v) triton X-100, 0.1% ampholytes, 50 mM DTT) applied to protect proteins. IEF gels were electrophoresed at 500V, 3mA/gel, 0.25W/gel, 7.5hrs for 6.5cm gels, and at 500V, 3mA/gel, 0.25W/gel, 17.5hrs for 15cm gels. After the run the gels were extruded using a water filled syringe into either 10%(v/v) glycerol for storage at -20°C or, transfer solution (0.3M Tris-base, 0.075M Tris-HCl, 3% SDS, 50 mM DTT, 0.01%(w/v) bromophenol blue) prior to loading on top of a 12% SDS-PAGE separating gel.

## 4.5.7 Barent & Elthon based 2D system

This protocol was adapted from the Barent & Elthon [1992] PAG-IEF with slight changes to the ampholyte composition. The gel mixture consisted of 335µl of stock acrylamide solution (28.4%(w/v) acrylamide, 1.6%(w/v) bis-acrylamide containing 5%(w/v) BioRad AF501-X8 ion exchange resin), 0.5ml stock triton X-100 (10%(w/v) triton X-100 containing 5%(w/v) BioRad AF501-X8 ion exchange resin) and 130µl of 40%(w/v) ampholytes (broad range 3-10 supplemented 45%(v/v) 5-8 ampholytes from Pharmacia) this was added to 1g of UltraPure urea. This mixture was placed into a water bath at 37°C to allow the urea to dissolve. Lyophilised samples were reconstituted in 20µl volumes of QH<sub>2</sub>O and 200µl of the gel mixture was added to each sample, 1µl of 10%(w/v) APS and 1µl of TEMED were then pipetted to initiate polymerisation. Immediately a 1.5mm i.d. x 5cm capillary tube was filled with this mixture by holding the tube horizontally and slowly dispensing the gel-protein mixture into the tube, polymerisation was usually complete within 30 minutes.

Capillary tubes were placed into the apparatus which contained anodic buffer (10mM phosphoric acid), the upper chamber was then filled with cathodic buffer (degassed 10mM sodium hydroxide). PAG-IEF was achieved by applying a constant 3mA/gel current (max 150V, max 1W) for 10 minutes switching to 300V for a further 10 minutes

then finally running at a constant 450V (max 3mA/gel, 1W max). Gels were extruded from the capillaries using a water filled syringe into 2x SDS-PAGE sample loading buffer and equilibrated for 5minutes. After equilibration they were then loaded onto a 12% SDS-PAGE separating gel and overlaid with 2x SDS-PAGE sample loading buffer, alternatively they were placed into 10%(v/v) glycerol for storage at  $-20^{\circ}$ C.

## 4.5.8 Ampholyte removal solution

A constant problem with free carrier ampholyte PAG-IEF systems is that the ampholytes migrate into the SDS-PAGE second dimension and stain deeply into a large smear that can obscure upto 20% of the gel. The ampholyte smear can be removed by incubating the PAG-IEF gel prior to equilibration or the second dimension SDS-PAGE gel after electrophoresis, in a solution of 47.5%(v/v) ethanol, 3%(v/v) acetic acid for >5 minutes.

## 4.6 Miscellaneous methods

A number of other methods were utilised during this research. The production of anti-TE antibodies was an attempt to show that ICF is truly of apoplastic origin and components do not appear in total extract (see section 5). The auxin affinity column methods were attempted towards the end of the project.

## 4.6.1 Production of rabbit anti-TE antibodies

In order to produce the maximum immune response from the rabbits a final protein concentration in the range 0.8 to 1.2 mg/ml is desirable. Tobacco TE was prepared as described in section 4.2.3 after 4 repeated VI's (see section 4.2.1) to remove soluble apoplastic proteins and polypeptides from the TE preparation. After ultracentrifugation the protein concentration was estimated using the BioRad protein estimation kit (section 4.3.2). A resultant concentration of 1.7 mg/ml of protein was calculated when read off a BSA standard curve (GraphPad Prism Software), i.e.: enough for injection. Antibodies were raised against both native proteins and denatured polypeptides to ensure that antibody probing could be carried out on native or denatured materials. Therefore the sample preparation included untreated native TE proteins and denatured TE proteins that had been boiled and contained 2% (w/v) SDS. An emulsion of this protein mixture was produced using the mineral oil, 'Freund's adjuvant' because it greatly enhances the immune response by prolonging immune system exposure to the antigens. This suspension was then injected into the rabbit; the injection regime was as follows: -

## Native-TE preparation for injections (- SDS)

- 1. 1.0ml of 1.7mg/ml TE containing 1mM DTT
- 2. Stored at -20°C until required for injection.
- 3. Thaw TE preparation.
- 4. 1.0ml of Freund's adjuvant; added in  $100\mu$ l aliquots, with vortexing and sonication.
- 5. Load 1ml into a 1ml syringe. Refreeze remaining millilitre for booster injection.

## Denatured-TE preparation for injections (+ SDS)

- 1. 1.0ml of 1.7mg/ml TE containing 1mM DTT
- 2.  $200\mu$ l of 10%(w/v) SDS (approximately 1.7mg/ml TE, 1mM DTT, 2%(w/v) SDS).
- 3. Place into a boiling water bath for 5 minutes.
- 4. Stored at -20°C until required for injection.
- 5. 1.0 ml of Freund's adjuvant; added in  $100\mu$ l aliquots, with vortexing and sonication.
- 6. Load 1ml into a 1ml syringe. Refreeze remaining millilitre for booster injection.

## Day 0 – Primary Injection

- 1. 0.5ml of +SDS administered intramuscularly into right leg thigh.
- 2. 10 x 50µl of +SDS administered subcutaneously along spine.
- 3. 0.5ml of -SDS administered intramuscularly into left leg thigh.
- 4. 10 x 50  $\mu$ l of -SDS administered subcutaneously along spine.

## Day 15 – Booster Injection

- 1. 0.5ml of +SDS administered intramuscularly into right leg thigh.
- 2. 10 x 50 $\mu$ l of +SDS administered subcutaneously along spine.
- 3. 0.5ml of -SDS administered intramuscularly into left leg thigh.
- 4. 10 x 50  $\mu$ l of -SDS administered subcutaneously along spine.

## 4.6.2 Generating and packing the auxin affinity column

Preparation of the epoxy-sepharose 6B was carried out as per manufacturers instructions (Pharmacia, Uppsala, Sweden). Briefly, 3g of sepharose was weighed into a 50ml Falcon tube and 20ml of  $QH_2O$  added. The sepharose was incubated for >30 minutes with agitation in order to re-swell the gel, (it swells to 4ml per original g of dried material). The sepharose was then filtered and washed for >1 hour with >600ml of  $QH_2O$  on a sintered glass funnel. It was calculated (see section 5.2.3.1) that 300µM concentration of hydroxyphenylacetic acid (HPAA) would adequate to saturate the

reactive oxirane groups contained within the gel. Consequently the gel was split into 2 equal weights and resuspended in the coupling solution (300 $\mu$ M HPAA) in a glass beaker and agitated at 45°C for > 24hours. After incubation the gel was washed extensively with >600ml of QH<sub>2</sub>O on a sintered glass funnel. Excess reactive groups were then blocked by incubating the gel in an excess of 1M triethanolamine >12 hours with agitation at 45°C. After blocking, the gel was washed with >600ml of QH<sub>2</sub>O and ligand coupling efficiency was assessed in one of two ways: -

*Indirect* – 536µl (approximately equivalent to 100mg of re-swollen sepharose) was centrifuged at 100g for 3 minutes. 400µl of supernatant was removed; the pellet was retained for the direct measurement. The absorbance of the supernatant was read against the original coupling buffer at the  $\lambda_{max}$  for HPAA of 277nm.

**Direct** – The pellet of sepharose from the indirect measurement (see above paragraph – 'Indirectly'), was washed by resuspension in 1ml of QH<sub>2</sub>O, centrifugation at 100g for 3minutes, and decanting of the supernatant this washing step was repeated 3 times. After washing the pellet was resuspended in 1ml of glycerol (the sepharose becomes optical clear) and the absorbance of the gel at  $\lambda_{max}$  for HPAA (277nm) was read against gel that had not been exposed to the coupling reagent and had been blocked with 1M triethanolamine.

The column was setup by utilising a peristaltic pump to the pack  $2 \times 5$ ml volumes of the auxin-sepharose into a  $2 \times 10$ ml syringe barrels at 133% of the maximum expected flow rate during usage – packing was therefore carried out at 1.33 ml/minute.

## 4.6.3 Running the auxin affinity column

The column was attached to a Pharmacia GradiFrac chromatography system at 4°C, equipped with a UV detector, fraction collector and a trace. The column was equilibrated with >4 column volumes (20ml) of 1M citrate-phosphate buffer, pH 5.5 containing 0.1M NaCI, at 1ml/minute, until a constant baseline was achieved at 280nm. Fresh ICF that had undergone dialysis for >24 hours against several changes of 1M citrate-phosphate buffer, pH 5.5 containing 0.1M NaCI was loaded (1-5ml volumes) at a flow rate of 0.3-0.5ml/minute. This sample was recollected and re-circulated through the column at 0.3-0.5ml/minute 3 times. The column was then washed with >300ml of 1M citrate-phosphate buffer, pH 5.5 containing 0.1M NaCI at 1ml/minute and elution of bound protein material carried out in one of two ways: -

*Elution Method No. 1* – was achieved by changing the buffer to 1M Tris-HCl, 0.5M NaCl, pH8.0 at a flow rate of 0.3-0.5ml/minute. The advantage of this method of elution

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was the absence of UV absorbing materials except eluting proteins consequently detection and subsequent collection of UV absorbing fractions is relatively simple.

*Elution Method No.* **2** – was achieved by changing to 1M citrate-phosphate buffer, pH 5.5 containing 0.1M NaCI and 10mM phenylacetic acid. This elution method was much more specific because the auxin (phenylacetic acid) specifically displaced any bound protein material from the column. Hence, one can report that eluted material can definitely bind to an auxin and it is therefore highly likely that the material is indeed an auxin-binding protein.

## 4.6.4 Cleaning and storing the auxin affinity column

The column was washed to remove any residual material with 100ml of 1M phosphate buffer at pH 8.0 containing 1M NaCl at 0.3-0.5ml/minute. For storage 50ml of 0.02% (w/v) sodium azide was passed through the column at 0.3-0.5ml/minute. The column was then sealed in parafilm and stored at  $4^{\circ}$ C.

## **SECTION 5**

## PROTEINS OF THE APOPLAST

## 5.1 Introduction

The technique of choice utilised throughout this research to extract the complement of proteins and enzymes present in the apoplast was vacuum infiltration (VI). This technique relies on the vacuum assisted infiltration of buffer into the apoplastic spaces to solubilise components followed by recovery of this intercellular fluid (ICF) with low speed centrifugation [Hammond-Kosack 1992]. This relatively simple technique is a powerful method to extract selectively and therefore partially purify apoplastic components primarily because the infiltration buffer is only in contact with the apoplast and therefore permits the selective extraction of apoplastic materials only. Ultimately, the major advantage of utilising a VI technique to extract apoplastic materials is that one can have a high degree of confidence that a particular component is actually present in the apoplast *in vivo*.

Contamination of ICF by intracellular components can and does occur and in order to confirm the apoplastic nature of materials extracted using VI it was common to use three 'assays' to confirm the apoplastic nature of the ICF. The first involved a visual assessment of ICF, if it was green it was discarded because intracellular contamination of ICF had undoubtedly occurred. The second was an assay for the intracellular enzyme, malate dehydrogenase (MDH) [Hammond-Kosack 1992], if the value contained in any ICF preparation exceeded 1% of the assay value of the total extract, it was deemed too high and was discarded [llett 1998]. The final assay was visual, and rested upon the fact that the large sub-unit of one of the most abundant leaf proteins, ribulose-1,5bisphosphate carboxylase (RUBISCO), [Chen et al. 2000] was readily identifiable as a prominent band at approximately 50 kDa on a SDS-PAGE gel. The presence of the RUBISCO large sub-unit in SDS-PAGE separated ICF preparations indicated that intracellular contamination had occurred and that any other polypeptides were not necessarily apoplastic in origin [Alexander 1995; llett 1998]. VI was the method of choice because other methods for the isolation of apoplastic/cell wall materials rely on the separation of these materials from whole cell and tissue extracts [eg: Shimuzi et al. 1997] and consequently will invariably contain higher levels of contamination from intracellular components than ICF.

In section 1.4 some of the wide range of proteins and enzymes contained within the apoplast that potentially can be selectively extracted using VI have been detailed. During

the course of this research two methods were employed to attempt to identify or attribute characteristics to VI extracted apoplastic proteins: -

- 1. N-terminal sequencing of polypeptides separated on SDS-PAGE gels
- 2. Spectrophotometric or in-gel enzyme assays (see section 7)

The ultimate aim was to catalogue and identify as many of the VI-extractable proteins and polypeptides as possible].

One group of proteins speculated to be present in the apoplast are the auxin-binding proteins (ABP's), see section 1.4.2. The apoplast is the proposed, but not yet clearly identified, site of action of ABP's. The highly selective nature of VI provided a very novel extract from which to attempt the isolation of ABP's. To the authors knowledge ICF has never been subjected to such a study and positive results would provide direct evidence of ABP presence in the apoplast *in vivo*.

## 5.2 Results

## 5.2.1 Efficacy, optimisation and development of VI techniques

The small scale vacuum infiltration (VI) technique previously used in Durham was initially trialed to ensure that it could provide the necessary selective enrichment of apoplastic polypeptides, this is detailed in section 5.2.1.1. The technique was then optimised to provide the largest quantity of apoplastic polypeptides with the lowest possible contamination from intracellular components (section 5.2.1.2). Section 5.2.1.3 depicts the data from VI's using buffers with differing properties in order to maximise the extraction of apoplastic proteins. Ultimately, any analysis of ICF extracted utilising the small scale VI technique was limited because of the relatively low quantities of material recovered, consequently, a large scale VI technique was developed which quadrupled the volumes of intercellular fluid (ICF) recovered (see section 5.2.1.4 and appendix 1).

## 5.2.1.1 The selective enrichment of apoplastic components

The first point within the small scale VI technique at which it was possible to make a qualitative assessment on the degree of intracellular contamination was immediately after centrifugation. Consequently, all recovered ICF's were subjected to a visual assessment of intracellular contamination based solely upon the colouration of the liquid. Leaf tissue was especially suited to such a 'quality assessment' because of the very high concentration of chlorophyll present in leaves, therefore any leakage of intracellular components (i.e.: the plasma-membrane and chloroplasts) during VI would impart a

green colour to the recovered ICF thus making a visual assessment of contamination relatively simple. Ultimately, any green colouration would bear a relationship to the degree of intracellular contamination present in the ICF and consequently, should improve the overall selective enrichment of apoplastic components achieved during VI.

To illustrate this point, three independent VI's were conducted on tobacco leaf tissue and the resultant ICF's were categorised as containing either LOW, MID or HIGH levels of intracellular contamination based purely on the visual assessment of the colouration of the liquid when viewed against a white background. Each category was then scanned over the visible spectrum, assayed for the intracellular enzyme malate dehydrogenase (MDH) from which the relative-MDH activity was calculated (section 4.3.1), the data are presented in figure 5.1. The scan revealed the presence of a peak at 682 nm in all ICF's which corresponded with the  $\lambda_{max}$  of chlorophyll a [Moberg *et al.* 2000] this was especially evident in the ICF visually assessed with a HIGH degree of intracellular contamination, but was also present to some degree in both MID and LOW ICF's. Similarly, the relative-MDH activity data (see the graph on figure 5.1) also reported the same trend of increasing intracellular contamination of ICF which 'matched' with the visual assessment and subsequent categorisation into LOW, MID and HIGH levels of contamination.

Polyacrylamide gel separation of ICF was the primary method of analysis to be utilised throughout this project, therefore equal protein loadings of the LOW, MID and HIGH visually assessed ICF samples were subjected to SDS-PAGE alongside tobacco leaf total extract (TE). Figure 5.2 illustrates the resultant gel. The TE lane contains a heavily stained band at ~50 kDa, this corresponds to the major sub-unit (the minor sub-unit is at ~15 kDa) of the most abundant tobacco leaf protein, RUBISCO [Chen at el. 2000], the band is arrowed on the figure. Due to the high abundance of RUBISCO in tobacco leaf any leakage of intracellular components during VI would be evident in ICF preparations because of the characteristic ~50 kDa polypeptide band. The conspicuous absence of RUBSICO in ICF preparations has been used previously to validate the apoplastic nature of ICF [llett 1998]. This polypeptide was definitely present in the HIGH ICF preparation and to a lesser degree in the MID ICF preparation. However, this band was not present in the ICF preparation that had been categorised as having a LOW level of intracellular contamination. All of these data therefore confirm that VI can selectively enrich components from the apoplast providing that these precautions for minimising intracellular contamination of ICF are undertaken: -

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Spectrophotometric and malate dehydrogenase enzyme assay of visually assessed intracellular contamination in ICF's.

3 vacuum infiltration's were carried out on tobacco leaf tissue using the small scale method (see section 4.2.1). The resultant ICF's were visually assessed for 'quality' (ie: lack of intracellular contamination) and categorised as having HIGH (obviously green in colour), MID (a hint of green colouration) or LOW (no obvious green colouration) levels of intracellular contamination dependent upon the degree of colouration. Each grouping was then pooled and scanned in the region 400-800nm in a Beckman DU7500 spectrophotometer (Beckman Instruments, UK) against infiltration buffer blank. A representative scan is shown top right. The same ICF's were also subjected to the MDH assay and the relative MDH-activity calculated as described in section 4.3.1, the results are presented on the graph, bottom right. Data are the means of ± SD of three measurements.





## SDS-PAGE analyses of ICF with varying degrees of intracellular contamination.

ICF was extracted from tobacco leaf tissue as described in section 4.2.1 and was categorised into either having LOW, MID or HIGH levels of intracellular contamination by visual assessment of colour. Each of the categories of ICF was dialysed overnight and then subjected to protein quantification (section 4.3.2), aliquoted into volumes that contained 50 $\mu$ g of protein and then lyophilised (section 4.2.4). The lyophilised solid was reconstituted in 50 $\mu$ l of SDS-PAGE sample buffer and subjected to SDS-PAGE as described in section 4.4.1. Key: M=denotes a 10  $\mu$ l loading of marker proteins; HIGH ICF=15 $\mu$ l (15 $\mu$ g) loading; MID ICF=15 $\mu$ l (15 $\mu$ g) loading; COW ICF=15 $\mu$ l (15 $\mu$ g) loading; TE=35 $\mu$ g loading. The gel was Coomassie stained (section 4.4.3).

- Visual assessment of ICF for green colouration prior to pooling,
- Ensuring that the relative MDH-activity is <1% of that found in TE, and,
- The major sub-unit of RUBISCO is not present in the ICF during gel analysis.

Previous work [Alexander 1995; Ilett 1998] had utilised anti-RUBISCO antibodies to establish the level of intracellular contamination present in ICF preparations. Unfortunately, these antibodies were not available for this study and as a consequence a different approach was taken. Native and denatured TE proteins were prepared from tissue that had been vacuum infiltrated four times in order to remove the majority of the soluble apoplastic protein & polypeptide complement as possible. This protein and polypeptide mixture was then injected into a rabbit to obtain an immune response (see section 4.6.1) the ultimate aim was to generate a spectrum of antibodies that had affinity for intracellular proteins only. TE and ICF proteins were subjected to SDS-PAGE and electroblotted for antibody probing as described in section 4.4.9. The optimal titre of primary and secondary antibodies was assessed by probing 16 individual TE blots with varying combinations of both antibodies, see figure 5.3; primary antibody titre number 2 and secondary antibody titre group B were determined to give the best banding patterns for subsequent analysis. Therefore, 1:1000 dilution of primary antibodies and 1:15,000 secondary antibody were used to probe an ICF blot (figure 5.3). Unfortunately, apoplastic proteins were also recognised by antibodies raised against TE proteins, presumably because the complete extraction of soluble apoplastic proteins may not been achieved or there was enough apoplastic protein material inside/outside the cells to cause a crossreaction. Ultimately these antibodies were not used again due to the high cross-reactivity.

## 5.2.1.2 Optimisation of the small scale VI conditions

It was noted that both the vacuum conditions and the developmental age of the leaf tissue played roles in the observed degree of infiltration; i.e., as the tissue was increasingly infiltrated with buffer it gradually becomes more, 'water-logged' and therefore darker in colour. A 'checker-board' experiment was set-up to optimise the observed degree of buffer infiltration into immature (top of plant, <10cm long) and mature (bottom of plant >20cm) tobacco leaf tissue exposed to the vacuum for varying lengths of time. Figure 5.4, illustrates the data from this experiment, and clearly shows that as exposure to vacuum increases so does the observed degree of buffer infiltration into the tissue. Furthermore, it was evident that immature tobacco leaves infiltrated to a greater extent than mature tobacco leaves. This presumably indicates that the apoplast of immature tissue has greater continuity for the conduction of liquids [Canny 1995]. As a



Anti-TE antibody probing of TE blots to assess optimum titre for probing ICF blot. Anti-TE antibodies were prepared as described in section 4.6.1. A total extract was prepared (section 4.2.3) and subjected to SDS-PAGE (section 4.4.1) and then blotted to nitrocellulose membrane (section 4.4.9) for antibody probing. Each number (1-5) represents a different dilution of primary rabbit anti-TE antibodies, as follows; number 1=1:100; number 2=1:1000; number 3=1:5000; number 4=1:10,000; number 5=1:25,000. Each group (A-C) represents a different dilution of secondary anti-rabbit conjugated alkaline phosphatase antibody (Sigma, UK), as follows; group A=1:5000; group B=1:15,000; group C=1:30,000. The ICF sample has been blotted and probed with a 1:1000 dilution of primary antibodies and a 1:15,000 dilution of secondary antibody.



## The effect of vacuum conditions on the infiltration of young and mature tobacco leaf tissue.

Several leaf strips were sectioned from immature and mature tobacco leaves and infiltrated with 100mM NaCl phosphate buffered to pH 7.5, vacuum conditions were: Exposure A = 15 sec. to evacuate ICF gas; 5 sec. agitation; 15 sec. repressurisation; Exposure B = 15 sec. to evacuate ICF gas; 15 sec. agitation; 15 sec. repressurisation; Exposure C = 15 sec. to evacuate ICF gas; 30 sec. agitation; 15 sec. repressurisation; Exposure C = 15 sec. to evacuate ICF gas; 30 sec. agitation; 15 sec. repressurisation; Exposure C = 15 sec. to evacuate ICF gas; 30 sec. agitation; 15 sec. repressurisation; Exposure D = 15 sec. to evacuate ICF gas; 60 sec. agitation; 15 sec. repressurisation; Control leaf strips are shown. Representative strips were scanned.

consequence of the inherently greater infiltration potential of immature tobacco leaf tissue it was used throughout this project, except where stated otherwise.

Two points were identified within the small scale VI technique that when optimised would maximise the yield of uncontaminated ICF: -

- In order to infiltrate the leaf tissue any gas present in the intercellular spaces must be removed. During the exposure to the vacuum the leaf tissue is in a depressurised environment and consequently cellular leakage may occur. Minimisation of the duration the tissue is exposed to a vacuum will reduce the level of intracellular contamination of ICF.
- 2. The recovery of infiltrated buffer from the leaf tissue requires the exposure of the leaf tissue to a centrifugal force; an excessive centrifugal force would cause the leaf tissue to be crushed at the bottom of the tube, and would thus directly elevate levels of intracellular contamination of ICF.

Therefore, using immature tobacco leaf tissue, the effects of the duration of exposure to the vacuum and the relative centrifugal force (RCF) used to recover the infiltrated buffer were investigated. The resultant measurements of the volume of visually uncontaminated ICF recovered, its protein content and the relative MDH-activities are graphed (A-C) on figure 5.5.

The volumes of visually uncontaminated ICF recovered at centrifugal forces of 500 and 700 were very low and when the tissue strips were examined they still appeared to be 'water logged' with buffer, it is therefore probably that the force exerted was not sufficient to drain the apoplast of infiltrated buffer. At 900 and 1200 g the tissue strips did not have the 'water logged' appearance, however, at 1200 g it was evident that some of the tissue was getting crushed at the bottom of the tube thereby reducing the volume of visually uncontaminated ICF recovered (figure 5.5, graph A). Consequently the maximum volumes of visually uncontaminated ICF were recovered after a 30 or 60 second exposure to vacuum and at a centrifugal force of 900 g. Similarly, at 900 g the protein content of ICF was relatively high (~280µg/ml) and because of the greater volume of visually uncontaminated ICF recovered at a 60 second vacuum exposure the overall yield was better. The final measurement of uncontaminated ICF is displayed on graph C, and shows that at 900 the ICF recovered after a 60 second vacuum exposure has a relative MDH-activity of ~0.8%, this corresponded to no visible RUBISCO on an SDS-PAGE gel (data not shown but see figure 5.2 for an example). The final optimised small scale VI technique is presented in section 4.2.1.



# Assessment of ICF's recovered from tissue using different conditions.

Duplicate VI's were carried out as described in section 4.2.1, except that the period of time that the leaf strips spent under vacuum was adjusted and the recovery of the ICF was undertaken at differing RCF's. After recovery, any ICF's that were visually contaminated were discarded and uncontaminated ICF's were pooled and the volume measured, the data is presented on graph [A]. Graph [B] - the protein content of each uncontaminated ICF was estimated using the Bradford assay (section 4.3.2). Graph [C] - the relative MDH-activity for each uncontaminated ICF was calculated as described in section 4.3.1. Data are the means of ± SD of two measurements.

## 5.2.1.3 Buffer selection

A series of vacuum infiltrations were conducted to select the most appropriate buffer to maximise the extraction of apoplastic materials while minimising any intracellular contamination of ICF. The buffers were chosen based on their different properties and previous use to selectively extract cell wall proteins from cell suspensions cultures [Robertson *et al.* 1997]. The buffers and their properties are presented in table 2.1.

VI Buffer	Extraction properties		
dH <sub>2</sub> O	Water soluble components only.		
50mM potassium phosphate, pH	Components that require low salt or neutral pH.		
7.5			
200mM sodium chloride, pH 7.5	Ionically bound components.		
200mM calcium chloride	Previously utilised for the extraction of cell wall		
	proteins.		
50mM EDTA	Cation sequestered proteins/polypeptides.		
50mM sodium borate, pH 7.5	Disruption of glycoprotein-polysaccharide		
	interactions.		

Table 2.1 – VI buffers	and their	properties	[Robertson	et al.	1997].
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Each buffer had very noticeable and individual effects on both the tissue and the resultant ICF. When infiltrated into the leaf tissue the calcium chloride VI buffer induced the loss of tissue 'elasticity' which was not regained after recovery of the infiltrated liquid, sodium chloride had a similar but lesser effect, this presumably stems from the altered properties of the cell wall in the presence of these ions [Greringer *et al.* 1995; Tibbits *et al.* 1998]. ICF recovered from tobacco leaf tissue was usually colourless or occasionally had a slight pink tint, however, ICF recovered after infiltration with the EDTA buffer had a deep pink-red colouration, no explanation of what components generate such a colour has been found. The spectrophotometric scan in figure 5.7 illustrates that the different ICF's recovered using buffers with different properties also exhibited different spectral properties.

Figure 576, A, shows the concentration of protein in each ICF (left Y-axis) and the final total quantity of protein recovered (right Y-axis). The sodium and calcium chloride VI buffers extracted the most protein, however, the calcium chloride ICF had a relatively high MDH-activity (graph B; the other buffers used all recovered ICF with relative MDH-activities of <1%) that presumably arose because the tissue was very difficult to handle after infiltration because of the loss of 'elasticity'. The ICF's were subjected to SDS-PAGE analysis to assess if intracellular contamination had occurred and to view the profile of



# Small scale VI using infiltration buffers with different properties.

Duplicate VI's were carried out as described in section 4.2.1 except that different infiltration buffers were used. Graph [A] - the concentration of protein in each visually uncontaminated ICF was assessed as described in section 4.3.2, the data is displayed on the left Y-axis; the volume of uncontaminated ICF was also recorded and used to the calculate the total quantity of protein recovered (right Y-axis). Graph [B] - the relative MDH-activity of each ICF was calculated against the appropriate TE prepared with the same buffer (see sections 4.2.3 and 4.3.1). Data are the means of ± SD of two measurements.



# Analysis of ICF's extracted using infiltration buffers with differing extraction properties.

See section 5.1 for detailed information. Duplicate VI's were carried out as described in section 4.2.1 except that different infiltration buffers were used. The contained 25µg of protein and then lyophilised (section 4.2.4). The lyophilised solid was reconstituted in 50µl of SDS-PAGE sample buffer and subjected to SDS-PAGE as described in section 4.4.1; each ICF lane was loaded with 10µg of protein. Key: 1=dH<sub>2</sub>O ICF; 2=PO<sub>4</sub><sup>3-</sup> ICF; 3=NaCl; 4=CaCl<sub>2</sub> ICF; 5=EDTA ICF's extracted with NaCI, CaCl<sub>2</sub> and borate infiltration buffers were spectrophotometrically scanned between 230 and 700nm against a water blank, the scan is shown on the left. Each of the ICF's was dialysed overnight and then subjected to protein quantification (section 4.3.2), aliquoted into volumes that ICF; 6=Borate ICF; 7=10µg loading of NaCI TE (see section 4.2.3) The gel was silver stained (section 4.4.4). extracted polypeptides. The gel image is displayed in figure 5.7. A low level of intracellular contamination (the RUBISCO polypeptide) is present in both the water and calcium chloride infiltrated ICF's, but is not present in the other samples. It is immediately obvious that the different buffers selectively extracted proteins and polypeptides from the apoplast. The 200mM sodium chloride buffer was chosen because of the combination of low intracellular contamination determined from the relative MDH-activity, the SDS-PAGE gel analysis, high total protein recovery and the similarity of banding patterns to the other VI buffers utilised (see section 4.2.1 for the full details of the final method).

## 5.2.1.4 Large scale VI

The small-scale VI method yielded relatively small volumes of ICF during a working day consequently a large scale VI technique was developed. The major difference was that bulk quantities of leaf tissue were rolled and therefore supported within a mesh during exposure to vacuum and centrifugation as a consequence the need to individually roll each tissue strip was removed. Therefore the time taken to VI comparable quantities of leaf tissue was greatly reduced and the volumes recovered greatly increased.

The technique was optimised in a similar manner to that of the small scale VI technique (see section 5.2.1.2). The data are not shown and described here because it is presented in a research paper that is currently undergoing submission to a peer reviewed journal. The paper is presented in appendix 1.

## 5.2.2 N-terminal sequence identification of ICF polypeptides

One of the main aims of this research project was to identify changes in the complement of apoplastic proteins. Therefore N-terminal sequence analysis was carried out on proteins present at different stages of leaf development (see section 5.2.2.1), during abiotic stress (see section 5.2.2.2) and finally, because of the substantial N-terminal blockage experienced, on as many polypeptides as could be clearly identified on a blot (see section 5.2.2.3).

## 5.2.2.1 N-terminal sequencing of ICF polypeptides during leaf development

Leaves were categorised as immature or mature as described in section 5.2.1.2, and then subjected to the fully optimised small scale VI technique (section 4.2.1). Similarly, age specific-TE's were prepared using immature and mature leaf tissues. All protein extracts were then subjected to SDS-PAGE separation. Gel lanes that contained marker proteins, ICF and TE were detached from each gel and Coomassie stained (see figure 5.8), the remainder of each gel was then electroblotted (section 4.4.11). The blots were



Band	M.W. (kDa)	Sequence	% homology	Description
ITL1	43.8	•	-	-
ITL2	33.9	-	-	-
ITL3	32.0	-	-	-
ITL4	26.0	ATFDIVNQDTYTVWA	86	thaumatin-like antifungal protein
ITL5	16.5	SVQDFCVADD	100	germin-like protein from arabidopsis
MTL1	63.8	······································	-	-
MTL2	62.4	-	-	-
MTL3	35.2	-	-	-
MTL4	31.5	•	-	-
MTL5	30.5	-	-	-

## N-terminal sequence analysis of immature and mature tobacco leaf polypeptides.

ICF was extracted from immature or mature tobacco leaf tissue as described in section 4.2.1. Proteins were loaded as follows; TE1=immature TE 35µg; TE2=mature TE 35µg; ICF1=immature 20µg; ICF2=immature 20µg. This gel was coomassie stained (section 4.4.3) and the duplicate was blotted (section 4.4.11) and selected polypeptide bands were then subjected to N-terminal sequencing. The table below the gel image details the results of sequence analysis and database searches for homology using BLAST (see section 4.4.12 for details of search criteria). Refer to the text for full information of the species and reference for each homology.

Coomassie stained and comparisons were made with the appropriate stained gel in order to select individual bands that appeared age-specific. Five immature (ITL) and five mature (MTL) polypeptide bands could be clearly identified on each respective blot and gel that appeared to be mutually exclusive of each developmental age, they were therefore subjected to N-terminal sequencing. The gel image and sequencing results are presented in figure 5.8 and the original BLAST search results are presented in appendix 2.

After several attempts using three different ICF preparations, sequencing revealed that 80% of the chosen polypeptides were seemingly either N-terminally blocked or composed of multiple bands. This level is high, but some studies have reported that in general plant proteins do tend to have a higher level of N-terminal blockage than other organisms. However, two of the ten polypeptides were successfully sequenced. A 29.2 kDa polypeptide which shared a significant (86%) homology with a 27 kDa thaumatin-like protein from Diospyros texana [Vu & Huynh 1994] and a 16.5 kDa polypeptide shared a 100% homology to germin-like protein from Arabidopsis thaliana.

## 5.2.2.2 N-terminal sequencing of ICF polypeptides during abiotic stress

There have been many reports of changes in the apoplastic protein complement in response to abiotic stress (see section 1.4.3.1 for an overview). Therefore tobacco plants were subjected to heat stress for 5 days at 40°C and drought stress was mimicked by restricting the water supply until all the leaves had wilted (~14 days) after which watering was resumed, the plants were then allowed to rehydrate over a 3 day period. Both sets of plants were then subjected to VI, SDS-PAGE and electroblotting.

A total of six polypeptide bands, which appeared to be induced by the imposed stress, were selected for N-terminal sequencing. Figure 5.9 displays the gel image and the selected bands. Several attempts were made but unfortunately, all of the selected polypeptides appeared to be N-terminally blocked or composed of multiple bands and therefore sequencing could not be carried out.

## 5.2.2.3 N-terminal sequencing of multiple ICF polypeptides from standard ICF

The consistent problem of N-terminal blockage of selected polypeptides had hindered the subsequent identification of apoplastic components. It was therefore decided that all of the bands that were clear, distinct and visible on a blot generated from standard ICF (see section 4.2.1) would be N-terminally sequenced. A total of 13 distinct polypeptide bands could be readily identified on the sequencing blot (SICF1-13) and the resultant gel image is presented in figure 5.10.



Band	M.W. (kDa)	Sequence	% homology	Description
HTL1	31.2	-	-	
HTL2	22.9	-	-	-
DTL1	50.2	-	-	-
DTL2	45.0	-	-	
DTL3	27.9	-	-	-
DTL4	19.4	-	-	-

## N-terminal sequence analysis of polypeptides from abiotically stressed plants.

ICF was extracted from heat stressed or drought stressed plants as described in sections 4.2.1 and 5.2.2.2. Proteins were loaded as follows; TE=NaCl TE 20 $\mu$ g; ICF1=heat treated ICF 20 $\mu$ g; ICF2=drought treated ICF 20 $\mu$ g. This gel was coomassie stained (section 4.4.3) and the duplicate was blotted (section 4.4.11) and selected polypeptide bands were then subjected to N-terminal sequencing. The table below the gel image details the results of sequence analysis and database searches for homology using BLAST (see section 4.4.12 for details of search criteria). Refer to the text for full information of the species and reference for each homology.



Band	M.W. (kDa)	Sequence	% homology	Description
SICF1	65.1	-	-	-
SICF2	46.9	MPVDITYLQSAVRKG RVPL	90	Putative pectinacetylesterase
SICF3	39.2	ASNGLGRTPQMGN	90	acetylgalactosaminidase
SICF4	38.2	-	-	-
SICF5	36	-	-	-
SICF6	34	-	-	-
SICF7	30.4	DDPD?T	-	1
SICF8	29.4	AVLDFCVGDLSVPDN PAG	77	Auxin-binding protein 20
SICF9	21.5	AVLDFC<	-	-
SICF10	20.3	•	-	-
SICF11	17.2	MPDP?PTFAE	0	No homology detected

## N-terminal sequence analysis of ICF polypeptides.

ICF was extracted from young tobacco leaves as described in sections 4.2.1. 40µg of ICF proteins and polypeptides were loaded to 5 duplicate lanes 4 were blotted and 1 was coomassie stained (shown above). This gel was coomassie stained (section 4.4.3) and the other lanes were blotted (section 4.4.11) and then the selected bands were subjected to N-terminal sequencing. The table below the gel image details the results of sequence analysis and database searches for homology using BLAST (see section 4.4.12 for details of search criteria). Refer to the text for full information of the species and reference for each homology. Key: ?=undetermined amino acid residue; <=sequencing reaction stopped.

Five polypeptide bands did not yield sequence data (~46%) either because of insufficient quantity of the polypeptide, the presence of more than one polypeptide or N-terminal blockage. Of the remainder only five polypeptides generated enough amino acid sequence to ensure a useful search of the database (>6 residues). Two homologies were present with carbohydrate related enzymes (SICF2 and 3) and two with an intercellular signalling protein (SICF8 and 9). The latter was an auxin-binding protein (ABP) isolated from prunus persica [Ohmiya *et al.* 1998]; both of these polypeptide bands had the same initial series of residues hence SICF9 sequencing was terminated by the operator of the sequencer because of the similarity to SICF8.

## 5.2.3 Attempted isolation of auxin binding proteins from ICF

Two ICF polypeptide sequences (SICF8 and 9) both yielded high homologies to an auxinbinding protein (ABP) identified in Prunus persica [Ohmiya *et al.* 1998]. Consequently, because the putative site of ABP action is assumed to be the apoplast [Brown & Jones 1994], ICF may well contain a disproportionate quantity of ABP's. Therefore, an affinity chromatography method was established in order to isolate and characterise any auxinbinding proteins present in ICF extracts.

## 5.2.3.1 Coupling hydroxyphenylacetic acid to epoxy-sepharose 6B

Several investigators had utilised hydroxyphenylacetic acid (HPAA) attached to a sepharose as an effective affinity matrix for the isolation of ABP's [Reinard & Jacobsen 1995; Kim *et al.* 1998]. The preparation of the epoxy-sepharose and subsequent HPAA-ligand coupling was carried out with reference to the manufacturers data and instructions (Pharmacia, Uppsala, Sweden) and as described in section 4.6.2. The following process was used to calculate the necessary concentration of HPAA for ligand coupling:-

Pharmacia quote that the epoxy-sepharose contains:-

19-40 $\mu$ M of reactive oxirane groups per ml of drained sepharose, and, 6-ml of sepharose in total was required for coupling to HPAA, therefore, a maximum total of 240 $\mu$ M oxirane groups for reaction with HPAA, the reaction between the oxirane group and HPAA is molar, therefore, a 300 $\mu$ M HPAA solution would provide enough ligand for coupling.

The efficiency of the coupling of the HPAA ligand to the epoxy-sepharose was assessed by following the decrease in absorbance of HPAA in the reaction mixture (figure 5.11, [A]) and by following the increase in absorbance of the optically clear epoxy-sepharose (figure



The auxin affinity matrix was prepared as detailed in section 4.6.2. These are the measurements of the coupling reaction (see section 5.2.3). Graph [A] = absorbance of the coupling mixture supernatant at different time points at the  $\lambda_{max}$  for HPAA of 277nm against dH2O. Graph [B] = absorbance of epoxy Auxin affinity column – graphs to show the progress and efficiency of the coupling of hydroxphenylacetic acid (HPAA) to epoxy-sepharose 6B. sepharose 6B at different time points at the  $\lambda_{
m max}$  for HPAA of 277nm against dH $_2$ O. Data are individual measurements. 5.11, [B]). The data are presented in figure 5.11 and illustrates that the coupling reaction was complete after approximately 24 hours.

## 5.2.3.2 Affinity chromatography of ICF

Any ABP's present in the apoplast were likely to be present at very low levels [Brown & Jones 1994]. Previous work has illustrated this point because of the need for highly sensitive microscopy techniques to visualise ABP's in the apoplast [Jones & Herman 1993; Deikmann *et al.* 1995]. In order to maximise the chances of binding ABP's to the auxin affinity column large volumes of ICF (5-ml) were freshly prepared and loaded at 0.5ml/minute. The ICF eluent was recollected and re-circulated through the column a total of three times. Elution of bound material was achieved either in a 'general' manner by altering the pH and ionic strength or more, 'specifically' by the addition of an auxin. The eluting fractions were collected, dialysed, lyophilised and subjected to SDS-PAGE, see figure 5.12 (A and B).

Elution using a change of pH and ionic strength (i.e.: 'general' - see 4.6.3) revealed that fraction 3 contained 6 polypeptides which had bound to the auxin affinity matrix, these are arrowed on figure 5.12, gel A. Of particular interest was a 21 kDa polypeptide (band F) and two larger polypeptides of 61 and 57 kDa (bands A and B, respectively), these polypeptides also eluted (fractions 3-5) with an auxin, indicating that they can specifically bind auxins (figure 5.12, gel B). The remaining polypeptides (C-E) were assumed to have bound to the sepharose matrix in a non-specific manner.

## 5.3 Conclusions

Early in this research it was proven that VI yields preparations of apoplastic material that are not visually or enzymatically contaminated by intracellular components when assessed by any of the methods to minimise contamination described within this thesis. Consequently, these methods adequately guard against intracellular contamination of ICF and lead the author to the conclusion that ICF extracted as described here is at the very least, a highly purified apoplastic extract.

During the development and optimisation of VI it was shown that polypeptides can be selectively removed from the apoplast using buffers with specified properties in a similar manner to that shown by Robertson *et al.* [1997] using cell suspension cultures. It was also evident that the different buffers had differing effects upon the tissue, in particular calcium ions induced the loss of elasticity. Powell *et al.* [1982] and Tibbits *et al.* [1998] have both shown that calcium ions can affect PGA's and thus alter the physical properties of the pectin matrix and Greringer *et al.* [1995] have illustrated that calcium ions can



## Analysis of bound fractions eluted from the auxin affinity column.

2x 5-ml volumes of ICF (see section 4.2.1) were recirculated through the auxin affinity column 3 times as described in section 4.6.3. Elution of bound material was achieved by changing the ionic concentration and buffer pH (gel A – elution method 1 in section 4.6.3) or specifically by supplementing the buffer with 10mM phenylacetic acid (gel B – elution method 2 in section 4.6.3). 1-ml fractions were collected, dialysed overnight against dH<sub>2</sub>O and then lyophilised (see section 4.2.4). Lyophilised material was reconstituted with 10 $\mu$ l of SDS-PAGE sample buffer, subjected to SDS-PAGE (section 4.4.1) and the resultant gels were then silver stained (section 4.4.4) for polypeptide visualisation. The arrows indicate silver stained polypeptide bands.

disrupt electrostatic interactions between cell wall polysaccharides that ultimately alter the extensibility of the cell wall. The phenomenon observed here is presumably the gross, overall effect of these ions.

N-terminal sequencing of apoplastic polypeptides was very disappointing because of the low yield of sequence data generated. Three potential reasons could explain this low vield. Firstly, it was possible, in some cases, that there was simply not enough of the chosen polypeptide present in order to sequence. As a result, several attempts were made to overload ICF extracts and therefore elevate the quantity of material present on the subsequent sequencing blot, however, this did not appear to improve the sequencibility of any of the chosen polypeptides. The second possibility was that many of the polypeptides selected for sequencing were in fact composed of many polypeptides. Thus it would be impossible to obtain a sequence. To counter this various acrylamide gel compositions were tried in order to effect a better separation of the chosen polypeptides, unfortunately none of these appeared to improve the subsequent sequencing attempts. Therefore, N-terminal blockage appeared to be the only possible mechanism by which so many of these polypeptides were resistant to sequencing. Consequently, methods were adopted to minimise any N-terminal blockage caused during SDS-PAGE (see section 4.4.11); these also did not alter the subsequent sequencibility of the polypeptides. Therefore the conclusion was drawn that many VI extractable apoplastic proteins are Nterminally blocked.

However, from the successful sequencing runs that were undertaken some very interesting and significant results were obtained. Immature tobacco leaf tissue yielded an apoplastic polypeptide of 16.5 kDa with a 100% sequence homology to a germin-like protein found in arabidopsis. Germin is a known extracellular protein of barley [Lane 1988] that actively produces hydrogen peroxide in tight correlation with tissue growth and is essential for cell wall maturation [Lane *et al.* 1992]. It is possible that a similar process may occur in the expanding tobacco leaf. Interestingly, this sequence also shares a 100% homology with a 19 kDa ABP isolated from peach [Ohimya *et al.* 1998] and although the protein did not exhibit hydrogen peroxide generating activity, wheat germin is also known to contain an auxin-responsive element [Lane 1994]. This raises the possibility of an auxin-mediated control of cell wall elongation during growth in tobacco leaf.

Immature ICF also contained a polypeptide with a mass of 26 kDa and good homology (86%) to a 27 kDa thaumatin-like antifungal protein isolated from Diospyros texana [Vu & Huynh 1994]. The presence of this protein in immature tobacco leaf tissue is unknown.

Unfortunately, abiotically stressed plants yielded only polypeptides with N-terminal blockages, which may reflect the stress the plant has been subjected to, i.e.: oxidative

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stress is a common phenomenon present in many types of stress (see section 1.5) and some oxidative species are known to induce the modification of the N-terminal region of animal proteins and polypeptides [Burcham & Kuham 1997] (see section 1.7). Coincidentally, the apoplast is also a known site of the production and release of oxidative species and this may explain why such a high level of N-terminal blockage was experienced during this study.

Standard ICF contained two polypeptides with polysaccharide and pectin related functioning, i.e.: they were extracellular matrix and cell wall related enzymes, not an unusual finding [Fry 1991]. Potential ABP's were again sequenced from ICF with molecular masses of 29.4 and 20.3 kDa; the latter corresponds to the molecular weight of an auxin-binding protein isolated from peach [Ohimya *et al.* 1998].

A surprising number of non-N-terminally-blocked proteins shared high homology with ABP's. Consequently, ICF preparations were subjected to chromatography through an auxin affinity matrix. Six proteins could be eluted from the affinity column with a simple change to the pH and ionic strength of the buffer. However, only three of these proteins were specifically eluted with an auxin supplemented washing buffer, thus indicating that these proteins were indeed *bona fide* ABP's. Unfortunately, there was not enough time to complete this study and obtain definite sequence confirmation that these proteins shared sequence homology with any known and characterised tobacco ABP's, but one of the putative ABP's was a 21 kDa polypeptide. A 20.3 kDa polypeptide was sequenced within this study (SICF9) that did share homology to an ABP, it is possible that these are true homologues of the same polypeptide and the slight difference in molecular mass can be accounted for by the day-to-day variation of the SDS-PAGE technique utilised in this study.

## **SECTION 6**

## THE DEVELOPMENT OF 2D ANALYSIS

## 6.1 Introduction

It was naively anticipated at the start of this part of the programme that 2-dimensional electrophoresis (2D) would be a relatively simple though highly resolving technique for the further separation of proteins and polypeptides observed on 1D gels (see section 5), in fact, the technique was deemed as an essential method of analysis primarily because it is the most powerful technique known for resolving complex protein mixtures [O'Farrell 1975; Ames & Nikaido 1976]. The resolution of a typical 2D-system is extremely high with the theoretical capability of separating individual polypeptides to homogeneity in one step, as a consequence small changes to the complement of the apoplast during stress and development could potentially be observed. For these reasons 2D analysis is rapidly becoming the method of choice for creating protein 'maps' or 'snapshots' of a particular tissue at a particular time point [Mollenkopf *et al.* 1999; Pardo *et al.* 1999; Yamaguchi & Pfeiffer 1999] and was therefore considered a useful tool for the study of the apoplast. A simple diagrammatic representation of a typical 2D-setup is presented in figure 6.1.

The most popular first dimension for 2D separations is polyacrylamide gel isoelectric focusing (PAG-IEF); this electrophoresis technique separates molecules based on their surface electrical charge. Two conditions must be met to effect an isoelectric separation of proteins and polypeptides: -

- 1. A pH gradient must be present in the gel, and,
- 2. The gel must not exhibit sieving properties.

The most common manner of establishing a pH gradient is the addition of 'carrier ampholytes' to a basic polyacrylamide gel mixture that contains no conducting species (i.e.: salts) and therefore has a very low ionic strength and poor ability to conduct current. Carrier ampholytes are comprised of hundreds of low molecular weight, polymeric and amphoteric species. When these compounds are exposed to an electrical field they migrate to their own isoelectric point (pl), at this position they are electrical neutral and therefore effectively do not migrate any further. Carrier ampholytes fulfil two crucial roles during isoelectric focusing, firstly they are very good conductors, and consequently are very effective at 'carrying' the current through the gel, secondly, they are very effective buffers, and thus they can establish a pH gradient once they have reached their isoelectric points.



## Figure 6.1

## The principle of 2-dimensional electrophoresis.

See section 6.1 for details. Briefly, complex solutions of proteins and polypeptides are applied to an isoelectric focusing (IEF) gel and electrophoresed until the proteins have reached the pl's and they are stationary. The IEF gel is physically transferred to a denaturing SDS-PAGE gel and electrophoresed to further separate the proteins and polypeptides on the basis of their molecular mass. The inset image (top right) is derived from Rüchel *et al.* [1978] and illustrates the pore formation characteristics of varying combinations of T (total acrylamides)/C (cross-linker) in polyacrylamide gels.

The second major pre-requisite for effective PAG-IEF is a polyacrylamide gel that does not appreciably sieve proteins and consequently does not hinder the movement of proteins and polypeptides through the gel matrix. Polyacrylamide gels are typically composed of a bis-acrylamide crosslinker (C) and acrylamide that is catalytically polymerised to form a porous matrix, the ratio of crosslinker to total acrylamides (T) determines the pore forming characteristics of the resultant polymerised gel. The inset figure (6.1) shows a collage of electron microscope images of polyacrylamide gels. In the horizontal series, C remains constant and as T increases left to right, the gel exhibits decreasing pore size and hence increasing sieving properties. On the vertical bar, T remains constant but C is varied, a parabolic relationship is observed which reflects the known sieving properties of polyacrylamide gels [Rüchel and Brager 1975; Rüchel *et al.* 1978]. Polyacrylamide gels are therefore practically non-sieving when gels are either dilute, the lowest concentration possible is 2%T, 2.2%C, or contain high levels of cross-linker, (%C >25).

The combination of non-sieving polyacrylamide gel containing a carrier ampholyte based pH gradient will cause proteins or polypeptides to migrate unhindered through the pH aradient while they still have surface charge and cease migration when they reach electrical neutrality, i.e.: their pl. Ultimately, the stationary zone the protein or polypeptides occupy at their pl will represent a dynamic equilibrium between pl-directed electrophoretic migration and diffusion away from the focused zone. A common problem encountered during PAG-IEF is the isoelectric precipitation of proteins and polypeptides at or near their pl; that is caused because they have little or no electrical charge and are present in an environment almost bereft of ionic species. As a consequence, PAG-IEF can be performed under native or denaturing conditions, typically, native PAG-IEF (contains no denaturants or detergents) is used only with proteins that are highly soluble throughout an isoelectric focusing procedure. Denaturing PAG-IEF, is the more commonly employed version of PAG-IEF because most proteins have a tendency to precipitate at their pl; typically, non-ionic denaturants and detergents such as, urea and triton X-100, are utilised to maintain the solubility of these proteins and polypeptides during the isoelectric focusing run.

The diagrammatic outline of 2D presented in figure 6.1 illustrates that after PAG-IEF, the gel is equilibrated in a denaturing buffer, usually SDS-PAGE sample buffer. This incubation permits sodium dodecyl sulphate to penetrate the gel and bind to the separated proteins and polypeptides, and allows the reducing agent (i.e.:  $\beta$ -mercaptoethanol or dithiothreitol) to break any disulphide linkages. Commonly, denaturation was aided by pre-warming the SDS-PAGE buffer to 70-80°C prior to submerging the PAG-IEF. After equilibration the PAG-IEF was then loaded onto the top of

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a SDS-PAGE and electrophoresed proteins and polypeptides that had been denatured in the equilibration step will now migrate into the SDS-PAGE gel and undergo separation based on their molecular mass. The highly resolving nature of this technique lies in the fact that proteins and polypeptides progress through two independent stages of separation; each stage separates based on completely different and individual physical properties of the proteins.

### 6.2 Results

### 6.2.1 Initial IEF trials

Effective use of 2D electrophoresis is highly dependent upon the reproducibility of banding patterns achieved during PAG-IEF, ultimately the spotting patterns observed on the resultant 2D 'protein map' must be superimposable between gels in order to accurately compare changes in the expression patterns of proteins. Consequently, a variety of PAG-IEF methods were evaluated early during this study to separate Pharmacia IEF marker proteins, test protein extracts, and ICF preparations. PAG-IEF gels of varying composition and formats were trailed, including, thin layer gels polymerised on microscope slides [Gorg et al. 1980], Pharmacia Multiphor immobilised pH gradient strips, a variety of horizontal and vertical slab and capillary tube systems [Tuszynski et al. 1979; Robertson et al. 1987; Baun et al. 1990]. All of these systems could effectively separate IEF markers and to some extent TE, but all invariably failed to effect an adequate separation of ICF's. One of the key problems experienced was that ICF's consistently formed a white precipitate after electrophoresis was started, which subsequently limited protein entry into the gel. This is a common phenomenon of certain protein mixtures (see sections 6.1 and 6.5) and has been traditionally circumvented by changing the sample preparation procedure prior to loading. Sample preparation procedures to remove contaminants that might induce precipitation [Granier 1988] and reduce IEF resolution were attempted but none succeeded effectively. Figure 6.2 shows the best PAG-IEF achieved during these trials; the technique used was an in-house adaptation of a vertical slab gel format system proposed by Robertson et al. [1987]. This gel was fairly representative of the PAG-IEF systems trialed, and indicated that ICF proteins do not enter the PAG-IEF; the poor entry of ICF-proteins and polypeptides in PAG-IEF gels was established as arising because of the rapid precipitation of ICFproteins that was visually evident very soon after the current had been switched on (<10 minutes).

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### Typical example of an analysis of ICF using PAG-IEF in a slab format.

BioRad Mini-Protean vertical gel 8 x 7cm format was used. The gels were cast at a thickness of 0.75mm from the following mixture: 7ml deionised water ( $QH_2O$ ), 2ml deionised acrylamide mixture (30%(w/v) acrylamide, 2% (w/v) bis-acrylamide), 2.4ml 50% glycerol and 0.6ml ampholytes (pH range 3-10). This mixture was degassed and polymerisation initiated by the addition of 50µl of 10% (w/v) ammonium persulphate and 20µl of TEMED, the gels were left to polymerise for 1 hour. The cathode solution was 25mM NaOH and the anode solution was 20mM acetic acid, both buffers were pre-chilled to 4°C prior to the electrophoresis. Samples and standards were prepared in a loading solution of final concentrations 30%(v/v) glycerol, 2%(v/v) ampholytes. Electrophoresis was conducted at a constant 200V for 1.5 hours followed by a further 1.5 hours at a constant 400V. After electrophoresis the gel was removed from the cassette and Coomassie stained (see section 4.4.3).

### 6.2.2 In-house adaptation of ESA Investigator™ 2D system

After several months of evaluation of many PAG-IEF systems without much success, the opportunity arose to visit the laboratory of ESA (Cambridge, UK), a commercial company specialising in 2D-electrophoresis systems. Separation of various age-related ICF's was achieved during this visit using the ESA-Investigator<sup>™</sup> 2D system – the final 2D-protein 'maps' obtained are exhibited in figure 6.3. The ICF material was extracted (as described in section 4.2.1) from tobacco leaves that had been graded into the three distinct leaf 'ages'; young (number 1), mid (number 2) or old (number 3). It was immediately evident that at least some ICF proteins did actually enter the PAG-IEF gel and were successfully separated by this 2D-system furthermore, these 2D gels did illustrate the potential of this highly resolving technique to discern changes in apoplastic proteins and polypeptides, for example: -

- Gp A, shows a triple spot pattern of proteins that does not seem to alter, at least qualitatively, as tobacco leaves age from young to old (figure 6.3 a-c).
- 2. B, illustrates a protein component that is either, lost or becomes unextractable from the apoplast as the tobacco leaf ages.
- 3. C, shows a new protein component apparently appearing during the mid-leaf age only.
- 4. A 'diffuse mass' that stains very heavily in the centre of all of the gels indicating that there is material in the sample that is present throughout leaf development that has heterogeneous charge spanning the pl range 5-8.
- >60 polypeptide spots were visible in the youngest leaf tissue and this number drops to <30 in the oldest ICF extracts (figures 6.3 a-c)</li>

However, even using this method it is evident that isoelectric focusing of many of the proteins and polypeptides had not reached the desired resolution; this was deduced because of the horizontal smearing present in the upper half of figure 6.3 (ICF 1) for example, and the general 'fuzzy' appearance of individual protein spots (see, 'B' or 'Gp A' on figure 6.3). This behaviour may be due to that fact that ICF material probably contains a range of glycoproteins with a wide degree of carbohydrate heterogeneity that would invariably lead to a wide degree of heterogeneity of isoelectric points as has been demonstrated previously for animal and plant derived glycoproteins [Dawes *et al.* 1994; Amoresano *et al.* 1999]. However, more disconcerting was that 400µg of ICF sample was required to produce each of the 2D gels presented in figure 6.3. The PAG-IEF sample preparation methodology used by ESA, included, acetone precipitation, denaturation with SDS and heat, and digestion of nucleic acids with nucleases (see section 4.5.2 for the full



# Analyses of ICF samples carried out in Cambridge using the ESA Investigator<sup>TM</sup> 2D electrophoresis system.

determination). Briefly; 400µg of each ICF was reconstituted, processed and loaded onto an IEF pre-cast immobilised pH gradient gel; sample preparation was as described in section 4.5.2 except using ESA pre-prepared reagents. After IEF the gels were transferred to gradient pre-cast Duracryl SDS-PAGE gels and mass separated. All gels have been stained with the ESA pre-prepared silver staining kit; GpA = is a group of 3 proteins that are present in ICF See section 6.3 for details. ICF's were recovered, dialysed and freeze-dried from leaves of different ages (see section 7.12 for details of leaf age throughout development; B= a protein that is apparently lost from ICF as leaves age; C= a new protein present as leaves age; Diffuse Mass = a dark staining area present throughout the leaf development series. procedure) and despite this a considerable quantity of precipitation was still evident when the electrophoresis procedure was started. Presumably this material was proteinaecous in origin because it had undergone such a rigorous sample preparation procedure for the extraction of proteins for application to the isoelectric focusing gel. Furthermore, 400µg of protein and polypeptide material separated into individual polypeptide spots in the final 2D gel should have been easily visible with only Coomassie staining, however, in all cases silver staining had to be adopted and as can be seen in figure 6.3 some of the gels are relatively over developed. It was apparent therefore that only a very small percentage of the total polypeptide material present in the ICF extract had actually entered the PAG-IEF gel.

The ESA method provided by far the best 2D separations of ICF material and thus it was decided to adapt equipment and techniques at Durham to develop an in-house version of the ESA Investigator<sup>™</sup> system. It was necessary to adapt equipment and techniques because the ESA Investigator<sup>™</sup> system cost in excess of £6,500, and resources for such a purchase were not available. Unfortunately, equipment and consumables of identical specification were difficult to acquire and it took over 1 month before a suitable power pack, capillary tubing and tube gel electrophoresis kit could be gathered together. Wherever possible identical equipment and parameters were used but certain modifications had to be made from the original ESA set-up; some of the more obvious included: -

- The dimensions of the capillary tubes used for PAG-IEF were different, consequently, voltage and current regimes had to be altered to effect similar separations.
- PAG-IEF gels used by ESA were precast using an immobilised pH gradient, importantly at Durham the PAG-IEF gels were cast using mobile carrier ampholytes, therefore consideration had to be given to the possibility of cathodic drift and increased potential for isoelectric precipitation.
- 3. Differences in power pack supply settings had to be considered.

All of these factors and others ensured that the adaptation of the methodology was not a direct replication of the ESA system. This meant that a number of trials varying electrophoretic duration, voltage, and current regimes based on extrapolated voltage, current densities derived from the ESA methodology and a variety of gel compositions, and formats were necessary to achieve a comparable separation of proteins. In order to monitor the PAG-IEF Pharmacia IEF marker proteins and a control protein mixture derived from mitochondria (see section 4.5.4 for preparation details) were utilised to ensure that pH gradient formation was successful. The control protein mixture was used

in preference to ICF during the method development for a couple of reasons, primarily because there was an abundant supply of material and the extraction of ICF was extremely time consuming and labour intensive, and secondly because, the opportunity existed to separate this material for a publication. Typically, to speed up the method development marker proteins and the control protein mixture were only subjected to PAG-IEF, after which proteins were visualised by staining the PAG-IEF. Furthermore a trial to assess the relative PAG-IEF separations in a small-scale format (6.5 cm) and a large-scale format (15 cm) were also tested.

Figures 6.4 and 6.5 illustrate the isoelectric separation of the marker proteins and control protein mixture, respectively under optimised conditions (see section 4.5.6 for in-house optimised electrophoresis conditions). Concomitantly, both the large and small format PAG-IEF capillary tube gels were analysed. Both of these formats yielded very similar separation patterns for both the marker proteins and the major components of the control protein mixture. Switching to the small format PAG-IEF system had the added advantage of greatly reduced electrofocusing times thus enabling higher sample throughput, in this case focusing in the small format system was reduced by 10 hours. The graphs on each figure show the Rf of each marker component in either the small or the large format system, it is obvious that the two formats essentially effect the same protein separation patterns.

Figures 6.6 and 6.7 illustrate that the hardware and conditions used in the ESA Investigator<sup>TM</sup> system were successfully adapted using in-house equipment and altered methodology, in both large and small scale formats. The pattern of proteins was extremely reproducible in both the large and small format systems, as illustrated by the comparison of protein separation in each (figures 6.6 and 6.7). The large format system does afford much better resolution but takes over 25 hours (including staining time) to complete a single gel, whereas, the loss of resolution observed in the small format system was to an extent compensated for by the greatly reduced running time, <10 hours (including staining time) for these particular gels. These images and the resulting antibody probed blots were presented at an international conference [see Memorandum, page v; Bowler *et al.* 1998].

Once again however, precipitation soon after electrophoresis had started was observed when attempting to run ICF in this adapted ESA system. Consequently, several variations of sample preparation were trialed (see sections 4.5.1, 4.5.2 and 4.5.3 for 2D sample preparation protocols), unfortunately, none yielded results because of the recurrent problem of polypeptide precipitation. Figure 6.8 is exemplary of the small format 2D gels obtained from ICF using this in-house adapted ESA system. The gel was greatly over stained to illustrate that very little protein or polypeptide material is actually present in the

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# IEF using marker proteins.

in section 4.5.6), were loaded with a 10µl aliquot of Pharmacia pl markers. After electrophoresis gels were 1x 15cm and 1x 6.5cm capillary tube PAG-IEF gels (composition and electrofocusing conditions are detailed extruded and stained with coomassie blue to visualise bands (see section 4.4.3). The relative mobility (Rf) was calculated for each protein and is plotted on the graph.





# IEF using control mitochondrial proteins.

1x 15cm and 1x 6.5cm capillary tube PAG-IEF gels (composition and electrofocusing conditions are detailed in section 4.5.6), were loaded with a  $50\mu$ l aliquot of control mitochondrial protein mixture (see section 4.5.4). After electrophoresis gels were extruded and stained with coomassie blue to visualise bands (see section 4.4.3). The relative mobility (Rf) was calculated for each major protein band and is plotted on the graph.



# 630µg loading of control mitochondrial proteins separated on large and small format 2D system.

100µl of a 42mg/ml solution of mitochondrial proteins (section 4.5.4) was prepared for IEF as described in section 4.5.2; the final pellet was resuspended in 100µl of IEF loading buffer. 15µl of this solution (630µg) was loaded onto a 15cm or a 6.5cm long IEF gel and focused for 17.5 and 7.5 hours, respectively (see section 4.5.6 for full protocol). After electrophoresis the IEF gels were extruded from their capillary tubes into SDS-PAGE (12%, 1.5mm thick) and electrophoresed. After electrophoresis the gels were silver stained (section 4.4.4). Note: the circle on each image relates to the sample buffer, equilibrated for 5 minutes and then loaded onto a large format SDS-PAGE gel (12%, 1.5mm thick) or a small format SDS-PAGE gel publication, Bowler et al. [1998]



# 630µg loading of heat shocked mitochondrial proteins separated on small and large format 2D system.

PAGE sample buffer, equilibrated for 5 minutes and then loaded onto a large format SDS-PAGE gel (12%, 1.5mm thick) or a small format SDS-PAGE gel (12%, 1.5mm thick) and electrophoresed. After electrophoresis the gels were silver stained (section 4.4.4). Note: the circle on each image relates to 100µl of a 42mg/ml solution of heat shocked mitochondrial proteins (see section 4.5.4) was prepared for IEF as described in section 4.5.2; the final pellet was resuspended in 100µl of IEF loading buffer. 15µl of this solution (630µg) was loaded onto a 15cm or a 6.5cm long IEF gel and focused for 17.5 and 7.5 hours, respectively (see section 4.5.6 for full protocol). After electrophoresis the IEF gels were extruded from their capillary tubes into SDSthe publication, Bowler et al. [1998]



### 250 $\mu g$ loading of tobacco leaf ICF

### Figure 6.8

### ICF proteins separated on a small format 2D system

 $250\mu$ g of freeze-dried ICF was reconstituted and processed according to ESA methodology (see section 4.5.2 and 6.3). The protein solution was applied to a 6.5cm long PAG-IEF gel in a  $20\mu$ l volume of IEF loading buffer. Focusing was 'achieved' by applying the voltage/current regime detailed in section 4.5.6 for 7.5hours, after which the gel was extruded into SDS-PAGE sample loading buffer and equilibrated for 5 minutes. The IEF gel was then loaded onto the top of a small format SDS-PAGE gel (12%, 1.5mm thick), electrophoresed. After electrophoresis the gels were silver stained (section 4.4.4). (For information, the cathodic end IEF is on the left-hand side of the slab gel).

second dimension gel. After several abortive weeks this methodology was abandoned because of the lack of progress.

### 6.2.3 In-house adaptation of the Barent & Elthon 2D system

During a survey of the literature a report of a rapid, small format capillary tube PAG-IEF and SDS-PAGE 2D system was found [Barent & Elthon 1992], this technique was adapted to the equipment available at Durham. Simultaneously the use of more subtle ways to follow the formation and maintenance of the pH gradient during PAG-IEF were used. Mixtures of coloured markers were developed which consisted of a combination of methyl red (pl~3), myoglobin (pl~7) and cytochrome C (pl~9.5) (see section 4.5.5 for preparation details). Figure 6.9 illustrates the use of such coloured markers during the optimisation of the Barent & Elthon adapted PAG-IEF method. The markers were invaluable, as it was much easier to view the location of a coloured marker in a glass capillary tube and thus actually 'view' the gradient forming. To confirm the visual data. capillary gels were routinely extruded and divided into 9 x 0.5cm segments, in triplicate, and placed into deionised water and after a set incubation period direct pH readings were taken. Figure 6.10 shows the pH readings displayed above each of the exhibited capillary gels. Figure 6.10 shows the same data plotted against time and confirms that the pH gradient in the system is rapidly lost after the first hour. The use of these two methods to view and monitor the formation of the pH gradient greatly sped up the trials of ampholyte composition, alteration of the anodic and cathodic buffer systems and changes to the duration of isoelectric focusing undertaken to optimise the system.

Figure 6.11 illustrates pH data generated as described above in the final optimised system (see section 4.5.7 for details of the final optimised method). Alteration to the range and ratio of ampholytes greatly reduced the pH gradient breakdown and ensured that the pH gradient is fully established within 1 hour and is maintained until 3 hours. The final isoelectric focusing run time was set at 2 hours and an example of mitochondrial control protein mixture run under these conditions is illustrated in gel A, figure 6.13. Separation was good and reproducible as judged by comparison with gel B; the whole 2D technique was extremely rapid. The gels in figure 6.12 were completed in less than 4.5 hours, including the 1-hour polymerisation time for the PAG-IEF gel. Gel A also exhibited a problem found throughout the development of all 2D methods that utilised mobile carrier ampholytes, 'ampholyte smear'. This is caused by the migration and intense staining of ampholytes with pl's>7 into the SDS-PAGE gel. Barent & Elthon [1992] also detailed an efficient ampholyte-removal step that can be applied to either the PAG-IEF gel prior to loading onto the SDS-PAGE gel or, to the SDS-PAGE gel prior to staining. Gel B figure 6.12 illustrates how the ampholytes can be selectively removed from a SDS-PAGE gel with a simple 5-minute wash with an, 'ampholyte removal solution' (see section 4.5.8) -

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### Alteration of the gel composition and ampholytes of the small format IEF and the effect on pH gradient formation.

Twelve 5cm IEF gels were loaded with  $15\mu$ l volumes of a mixture of coloured pl markers, see section 4.5.5 for preparation details. IEF was conducted as described in the legend that accompanies figure 6.10, but the gel composition and buffer system were adjusted (see section 6.2.3 for details). At the specified time-point electrophoresis was stopped and each group of 3 gels were then extruded, scanned, divided into 9 pieces, each 0.55cm in length, placed into 1.5-ml of QH<sub>2</sub>O and incubated for 2hours. After incubation the gel piece was removed and the pH of each of the resultant solutions was measured, the data is presented on the graphs above.



### Removal of the ampholyte 'smear'.

Gel A =  $300\mu$ g of control mitochondrial proteins was electrophoresed in the following sequence: 10minutes at 100V max.; 10minutes at 250V; 100minutes at 400V through a 5cm IEF gel using the altered electrode buffer system and gel composition (see section 6.4) and then subjected to small format SDS-PAGE (12%, 1.5mm thick). Gel B =  $200\mu$ g of control mitochondrial proteins was electrophoresed in the following sequence: 10minutes at 100V max.; 10minutes at 250V; 100minutes at 400V through a 5cm IEF gel using the altered electrode buffer system and gel composition (see section 6.4) and then subjected to small format SDS-PAGE (12%, 1.5mm thick). Gel B =  $200\mu$ g of control mitochondrial proteins was electrophoresed in the following sequence: 10minutes at 100V max.; 10minutes at 250V; 100minutes at 400V through a 5cm IEF gel using the altered electrode buffer system and gel composition (see section 6.4) and then subjected to small format SDS-PAGE (12%, 1.5mm thick). Gel A was immediately Coomassie stained (section 4.4.3); Gel B was treated with the, 'ampholyte removal solution' (see section 4.5.8 and 6.4) and then Coomassie stained in the usual manner (see section 4.4.3).

the resultant stained gel of control mitochondrial protein mixture revealed an additional number of previously obscured proteins.

Several attempts were made to separate ICF using this methodology (section 4.5.7) preparing the sample using the intricate ESA acetone based sample preparation methodology that had given reasonable results in section 6.3 (see section 4.5.2). Unfortunately, precipitate was obvious and no 2-D pattern was produced, see figure 6.13, gel A. A variety of other sample preparation methods were attempted (see sections 4.5.1 and 4.5.3 for 2-D sample preparation protocols), but again all failed apparently due to excessive precipitation of proteins, figure 6.13, gels B & C show examples.

A new loading strategy was devised after consultation with the authors of the Barent & Elthon paper. The strategy was unusual in that the sample was incorporated directly in the gel mixture prior to polymerisation. This strategy worked exceptionally well for the control protein mixture and enabled the isoelectric focusing of up to 500µg of protein material. Also this strategy did yield a protein pattern for ICF, see figure 6.14 – but not what was expected. It was noticeable that ICF protein or other components loaded in this manner still precipitated out in the gel during electrophoresis and effectively turned the gel white.

### 6.3 Conclusions

A significant number of IEF systems were tried initially, but it was found to be extremely difficult to get a PAG-IEF system to work reproducible when separating ICF material. Throughout the method development of 2D undertaken during this research project the ability to effectively monitor the formation of the pH gradient during PAG-IEF had been the limiting step. Consequently, methods to directly monitor the gradient were devised and rapidly speeded up progress towards the eventual development of two 2D systems each with their own unique merits. The ESA adapted system yielded excellent 2D separations of mitochondrial proteins good enough to be presented at an international conference [see memordandum; Bowler *et al.* 1998]. Furthermore, scaling down the procedure proved useful for the development of a rapid throughput 2D system that could yield protein 'maps' within 1 day. The second 2D system that was developed was based upon a similar small scale system described by Barent & Elthon [1992] and after trials a unique in-gel loading strategy was undertaken that could separate upto 500µg of polypeptide material in a single gel (data not shown). This potentially opens up the possibility of traditional N-terminal sequencing from 2D blots.

Unfortunately, good reproducible separation of ICF material was never achieved within the time constraints of the project primarily because of the constant problems with



### Separation of ICF using an in-gel loading strategy.

 $150\mu$ g of freeze-dried ICF was reconstituted in IEF gel mixture; the gel/sample mixture was briefly microfuged to pellet any precipitate and the supernatent collected. The gel was polymerised by the addition of ammonium persulphate. The normal 2-D separation protocol was followed (see sections 4.5.8 and 6.4) and the resulting gel was Coomassie stained (see section 4.4.3).

precipitation of ICF components. However, even the best separations achieved using the ESA Investigator <sup>™</sup> 2D-system exhibited quite marked precipitation, horizontal smearing and 'fuzzy' protein spotting patterns, this can presumably be explained by the heterogeneous nature of the glycoprotein materials extracted from the apoplast and/or the presence of potentially heterogeneous contaminants such as phenolic compounds and carbohydrate material [Fry 1991; Dawes *et al.* 1994; Dietz *et al.* 1997]. The constant and rapid precipitation of ICF in an isoelectric focusing system is evidently a feature of plant apoplastic proteins and polypeptides and presumably belies the ionically rich environment that this material normal resides in, isolation into an ionically poor buffer coupled with huge heterogeneity in charge may go someway to explain the precipitous behaviour of these proteins during PAG-IEF.

### **SECTION 7**

### APOPLASTIC ANTI-AOS ENZYMES

### 7.1 Introduction

Active oxygen species (AOS) include the superoxide anion ( $O_2$ ), hydrogen peroxide ( $H_2O_2$ ) and the hydroxyl radical (OH). These molecules are the highly reactive and consequently very deleterious by-products of aerobic respiration and photosynthesis [Richter 1995]; see section 1.5 for more detail. The intracellular environment contains anti-AOS defences (see table 1.6) that can absorb the oxidative potential of AOS (ubiquinol, vitamin E, etc) and a range of enzymes that can catalytically convert AOS (superoxide dismutase, etc), these defences provide effective protection against intracellular AOS.

It may therefore be surprising to find, that AOS are also biologically useful molecules that are deliberately generated and released into the apoplast during particular developmental stages in order to oxidatively couple cell wall polymers and proteins [Canny 1995; Dietz 1997] and in response to pathogen attack because of their anti microbial activity [Chen *et al.* 1994; Bergmann *et al.* 1994; Broekaert *et al.* 1995; Brisson *et al.* 1994]. Surprisingly, abiotic stress such as, temperature [Marentes *et al.* 1993], wounding [Bradley *et al.* 1992; Bergmann *et al.* 1994], drought [Covarrubias *et al.* 1995], and toxic chemicals [Brune *et al.* 1994; Blinda *et al.* 1997] are all known to elevate the levels of AOS, a similar elevation has also been observed during ageing and senescence of plant tissue [Rogiers *et al.* 1998]. The exact mechanisms for abiotic- and age-related elevation of AOS have not yet been elucidated.

The observable presence of AOS in the leaf apoplast is proposed to occur via one or a combination of three independent processes (see figure 1.6 and section 1.5 for an overview): -

- The action of an NADPH-oxidase complex regulated from the cytosol [Sutherland 1991; Vera-Estrella *et al.* 1992; Scheel 1998].
- The action of apoplastic and cell wall peroxidases and oxidases [Lane 1994; Wojataszek *et al.* 1997; Fry *et al.* 1998; Schweikert *et al.* 2000].
- 3. A general age-related decline in anti-AOS defences that would ordinarily remove any AOS present [Rogiers *et al.* 1998].

In addition to being the site of AOS accumulation the apoplast also contains a range of anti-AOS defences presumably to manage levels of AOS; these include antioxidant compounds, such as, glutathione, ascorbate, polysaccharides and phenolic compounds and a suite of enzymes that can directly metabolise AOS's such as catalases, peroxidases and superoxide dismutases (see table 1.6 and section 1.5). Unfortunately, relatively little information is known about the occurrence and action of apoplastic anti-AOS enzymes and in fact until very recently many had been considered to be purely intracellular, for example, superoxide dismutase [McCord & Fridovich 1969; Vanacker *et al.* 1998]. As a consequence, of the scant literature concerning the anti-AOS enzymes present in the apoplast it was a relatively logical extension of this research project to further investigate these enzymes.

The superoxide dismutases (SOD's) are the only known enzymes that can dismutate the superoxide anion, therefore reducing the potential for the generation of the more deleterious, hydroxyl radical [McCord & Fridovich 1969]. All SOD's have been characterised into one of three groups depending upon which metal ion they co-ordinate. The most primitive form of SOD contains an iron ion (Fe-SOD) or a manganese ion (Mn-SOD), and are typically found in bacteria, algae, mitochondria and therefore, of course, in plant and animal cells. More recently evolved forms of SOD containing the metal ions, copper and zinc (CuZn-SOD) have been located to the chloroplast and cytosol [reviewed in Bannister *et al.* 1987]. SOD was, until relatively recently, considered to be purely an intracellular matrix [Strålin *et al.* 1995] and the recent localisation of CuZn-SOD to the plant apoplast [Streller & Wingsle 1994]. This new information has lead to a re-evaluation of the roles of SOD's in both animals [Strålin *et al.* 1995] and plants [Streller & Wingsle 1994].

Peroxidases (POD's) are ubiquitous throughout the primary cell walls of all higher plants and are often expressed in a tissue-specific, developmental- and stress-related manner. Therefore, they are frequently implicated in differentiation, pathogen resistance and wound response [Rao *et al.* 1990; Cordewener *et al.* 1991]. POD enzymatic action is integral to apoplastic integrity, development and intercellular signalling [Mader 1980; Fry 1986]. It is also clear that apoplastic POD activity rises in response to virtually all abiotic and biotic stress [Dietz 1997] indicating major roles for POD in stress responses and anti-AOS defence. POD's can be classified as either cyanide-sensitive (Fe-POD's) or insensitive (Se-POD's) depending upon their reaction to the cyanide ion [Rob *et al.* 1997]. The primary aim in this part of the study was to positively and unequivocally identify apoplastic SOD's, POD's and catalases, preferably by N-terminal sequencing and homology searching. A range of convenient, native-PAGE in-gel assays, specific for each of the anti-AOS enzyme groups of interest was utilised to locate bands with activity. These methods were a rapid way to partially purify and simultaneously identify components within relatively complex apoplastic extracts with anti-AOS enzyme activity. Once bands of activity had been identified, characterisation was attempted using SDS-PAGE, enzyme inhibition studies and N-terminal sequencing.

### 7.1.2 Notes on Experimental Design

Figures 7.1, 7.2 and 7.6 illustrate enzyme activity staining in native-PAGE gels for the enzymes catalase, peroxidase (POD) and superoxide dismutase (SOD), respectively. Materials for the gel analyses were prepared in native sample buffer and extracts applied to the wells of native-PAGE gels (see section 4.4.2), after which the gel was removed from the cassette and submerged in the appropriate staining solution until stained to the desired intensity (see sections 4.4.5-7). Gels were then scanned using a flat bed scanner attached to a PC. Initial trials with SOD and POD stains revealed that the two procedures could be applied to a single gel providing that the gel was washed for >3 minutes in pure water between staining procedures.

Three independent physical properties, electrical charge, overall mass, and the shape of the native protein dictate how they migrate in such an electrophoresis system. As such, it is difficult to assign unequivocally physical characteristics to individual bands separated in native-PAGE systems; however, it is possible to make broad and simplistic generalisations. Small proteins with high pl's will tend to migrate further down the gel towards the cathode, while on the other hand, larger proteins with more acidic pl's will either not enter the gel or will not progress very far.

In sections 7.8 and 7.9 attempts were made to assign molecular weights to components that had specific enzyme activities. A number of trial attempts were made to elucidate the most effective way of removing an enzymatically active band from a native-PAGE gel and to transfer it to denaturing system (data not shown) in order to perform SDS-PAGE and subsequent mass estimation. Eventually a similar method to that of 2-dimensional electrophoresis was chosen; bands were excised and incubated for 5 minutes in warmed denaturing SDS-PAGE sample buffer, loaded on the top of a denaturing separating gel, overlaid with more SDS-PAGE sample buffer and then electrophoresed.

Comparison of in gel POD and SOD enzyme assays of TE and ICF samples (i.e.: figures 7.2 and 7.5, but also throughout section 7), illustrated either a potential overlap between the anti-AOS enzymes present in the apoplast and the intracellular environment or a degree of contamination. Both are possible explanations, but it should be noted that VI is not a perfect separation technique for the isolation of pure and uncontaminated apoplastic protein material and it will always have the potential to contain some degree of intracellular contamination. What can be said is that VI, although not ideal, does selectively and preferentially concentrate apoplastic material in relation to intracellular contaminants. As such, subsequent in gel enzyme activity analysis should be viewed as a qualitative assessment of the existence of unique extracellular POD's and SOD's and quantitative judgement should only be made with great care.

### 7.2 Results

### 7.2.1 Attempted detection of apoplastic catalase

Figure 7.1 (gel A) illustrates that control enzyme (commercially available catalase -Sigma, UK) can be detected as a yellow to pale blue band on a dark blue background at the bottom of a native gel. Unfortunately, catalase from tobacco leaf TE's could not be identified on the same gel. This was assumed to be due to the extremely dark blue background staining that may have masked any pale blue to yellow bands of enzyme activity. Consequently, the concentration of the enzyme substrate, hydrogen peroxide, was reduced from 3% (v/v) to 0.3% (v/v) in order to decrease the level of dark blue background staining, the result is illustrated in figure 7.1, gel B.

Gel B shows that alteration of the concentration of the substrate did improve the overall clarity of the detection system, as indicated by the improved resolution of the control enzyme, but still no catalase could be readily identified in TE or ICF preparations. A number of possibilities can be thought of to explain this negative result; catalase may have never entered the gel; it is possible that catalase has strong ionic interactions with the plant cell wall and the infiltration buffer was not of sufficient ionic strength to break the interactions; it is also possible, that the sample preparation procedure of extensive dialysis and lyophilisation had inactivated any catalase enzyme present in the ICF. Therefore, this assay was not developed further within this study.

### 7.2.2 Detection and optimisation of separation of apoplastic POD's

Staining for POD enzymes capable of utilising the artificial POD substrate guaiacol was much more fruitful (see figure 7.2, gels A and B). The control enzyme (commercially



Assay of catalase enzyme activity of control catalase enzyme, TE and ICF in native-PAGE gels.

TE and ICF were prepared as described in sections 4.2.3 and 4.2.1, respectively. Both gels were 10% (w/v) native-PAGE gels and samples were prepared in native PAGE sample buffer (see section 4.4.2). The quantities quoted refer to either Bradford quantified protein concentration or quantities of protein/enzyme activity was derived from the information from the commercial supplier (control POD enzyme – Sigma, UK). Gel A was loaded as follows; TE =  $35\mu g$  of total extract; CAT1 =  $89.55 U (45\mu g)$  of control enzyme; CAT2 =  $29.85 U (15\mu g)$  of control enzyme; and assayed at a substrate concentration of 3% (v/v). Gel B was loaded as follows; ICF =  $35\mu g$  of sodium chloride extract ICF; CAT =  $17.91 U (9\mu g)$  of control enzyme; TE =  $35\mu g$  of total extract; and assayed at a substrate concentration of 0.3% (v/v). See section 4.4.5 for a detailed description of the catalase staining technique.



Assay of peroxidase enzyme activity of control peroxidase enzyme, TE and ICF in native-PAGE gels.

TE and ICF were prepared as described in sections 4.2.3 and 4.2.1, respectively. Both gels were 10% native PAGE gels and samples were prepared in native PAGE sample buffer (see section 4.4.2). The quantities quoted refer to either Bradford quantified protein concentration or quantity of protein/enzyme activity was derived from the information from the commercial suppliers (control POD enzyme – Sigma, UK). Gel A is loaded as follows; POD1 =  $1.8 U (30\mu g)$  of control enzyme; POD2 =  $5.4 U (90\mu g)$  of control enzyme; TE =  $17.5\mu g$  of total extract. Gel B is loaded as follows; TE =  $35\mu g$  of total extract; POD =  $0.9 U (15\mu g)$  of control enzyme; ICF =  $35\mu g$  of sodium chloride extract ICF. See section 4.4.6 for a detailed description of the POD staining technique.

available enzyme - Sigma, UK) migrated to the upper third of the gel and was easily visible as two red-brown densely staining bands on a clear background.

TE protein from tobacco leaf yielded two well-defined bands that migrated to the middle of the gel (TE-POD 1 & 2). The same loading of ICF protein also indicated that these enzymes were present in the apoplastic preparation, but in addition 4 further bands of POD enzyme activity were present, 2 above and 2 below what had been designated as TE-POD 1 & 2 (see figure 7.2, gel B). POD enzyme active bands located only in ICF were named ICF-POD 1-4 progressing from the top of the gel to the bottom.

In order to qualify the assumption that the four identified ICF-POD's were truly apoplastic the pattern of POD enzyme staining bands in TE was examined using a higher loading range (5-150µg) of protein, (see figure 7.3, gel A). Analysis of the TE-POD enzyme patterns illustrates that TE-POD's 1 and 2 appear to be the only peroxidase active bands in TE. Furthermore, even when 150µg of TE protein was loaded the unique POD enzyme bands, designated ICF-POD's 1-4 that have only been observed in ICF were not detected in TE preparations. It was concluded that ICF-POD's 1-4 are apoplastic in origin because they are unique to ICF samples as determined by POD activity staining.

Figure 7.3, gel B, illustrates a gel with increasing protein loadings of ICF and clearly shows that ICF-POD's 1 and 2 and ICF-POD's 3 and 4 are grouped into two pairs. In an attempt to achieve a better separation of these proteins and to reveal if the four bands of enzyme activity present in the ICF preparations were composed of more components, the same ICF preparation was subjected to native-PAGE electrophoresis in gels of varying acrylamide concentrations (see figure 7.4). The 10% (w/v) acrylamide native-PAGE gel clearly demonstrated that the upper ICF-POD's was definitely comprised of two distinct bands of POD enzyme of activity. ICF-POD's 3 and 4 did not separate as effectively but remained as two distinct bands of POD enzyme activity (see figure 7.4, 10% and 12.5% (w/v) gels).

### 7.2.3 Spectrophotometric assay of POD activity in excised bands

In order to re-confirm that ICF-PODs 1, 2, 3 and 4 actually contained POD enzyme activity, two groups of duplicate bands were carefully excised from a POD stained native gel and placed into eppendorfs.  $750\mu$ I of 100mM phosphate buffer, pH 7.0 was added and the gel was macerated for >1 minute. The buffer and gel mixture was incubated for 24 hours at 4°C, after which the gel was pelleted in a microfuge and the supernatants collected and assayed for POD activity spectrophotometrically as described in section 4.3.4 (using a Beckman DU7500 spectrophotometer). The activity data derived from



### Assay of peroxidase activity in TE and ICF separated in native-PAGE gels.

TE and ICF were prepared as described in sections 4.2.3 and 4.2.1, respectively. Both gels were 10% native PAGE gels and samples were prepared in native PAGE sample buffer (see section 4.4.2). The quantities quoted refer to Bradford quantified protein concentration. See section 4.4.6 for a detailed description of the POD staining technique. Gel A has initially been stained for SOD activity and is loaded left to right, as follows; Lane 1 =  $5\mu$ g TE; Lane 2 =  $20\mu$ g TE; Lane 3 =  $35\mu$ g TE; Lane 4 =  $50\mu$ g TE; Lane 5 =  $75\mu$ g TE; Lane 6 =  $100\mu$ g TE; Lane 7 =  $125\mu$ g TE; Lane 8 =  $150\mu$ g TE. Gel B is loaded left to right, as follows; Lane 1 =  $17.5\mu$ g ICF; Lane 2 =  $35\mu$ g ICF; Lane 3 =  $52.5\mu$ g ICF.



## Assay for peroxidase enzyme activity in ICF separated in native-PAGE gels of differing compositions.

All gels are native (see section 4.4.2) but of differing acrylamide ratios (the actual acrylamide percentage (w/v) of each gel is indicated above the gel image). All gels were loaded with 35µg of Bradford assay quantified protein from the same sodium chloride extracted ICF. All gels were initially stained for SOD enzyme activity and then counterstained for POD enzyme activity. See section 4.4.6 for detailed descriptions of the staining techniques.

duplicate measurements were expressed as percentages of the total POD activity present in the same quantity of protein. Unfortunately, measurement of the recovery of protein from the gel extract was not attempted, and the potential for enzyme inactivation had not been considered consequently, direct comparisons cannot be made. However, it appears that approximately 26% of the total POD enzyme activity present in ICF is contained within ICF-POD's 1-4 as separated by native-PAGE even after electrophoresis and a 24 hour incubation period (see figure 7.5).

### 7.2.4 Detection and optimisation of separation of apoplastic SOD's

Using a SOD-specific native in-gel staining assay (see section 4.4.7), it was possible to identify the SOD control enzyme (commercially available SOD – Sigma, UK) on native gels as a large clear zone on a dark blue background (see figure 7.6, lanes SOD1 and SOD2) in the top third of the gel.

TE extracted from tobacco leaf tissue exhibited 4 distinct bands of SOD enzyme activity in the middle section of the native gel, these bands were named TE-SOD's 1-4, see figure 7.6, gels A and B; it was also noticeable that TE exhibited a high background level of SOD enzyme activity that presented as a smeared area in the upper portion of the native gel. Unfortunately, this SOD enzyme component could not be directly attributed to one or more specific bands of activity thus revealing that some proteins are not resolving well in this native-PAGE system.

On a protein weight-for-weight basis gel B (figure 7.6) clearly shows that ICF also contains an area of high SOD activity on the upper portion of the gel, this area is much more active than observed in TE, and potentially indicated that vacuum infiltration has selectively enriched a true apoplastic SOD. ICF also exhibits 2 of the 4 bands of SOD activity present in TE (TE-SOD's 1 & 2) in the middle section of the native gel, this may arise because of some degree of intracellular contamination of ICF has occurred or because these particular enzymes are present in both compartments. In the lower half of the native gel distinct groupings of SOD-active bands are visible that are not present in TE.

In order to ensure that the bands that had stained enzymatically positive for ICF-SOD activity were not an artifact of a low TE loading, a range of  $5\mu$ g to  $150\mu$ g of TE was loaded and run through a 10% (w/v) native gel (see figure 7.7, gel A). A similar loading study was performed with ICF with a view to enhancing the distinct bands present in the lower portion of the native gel (see figure 7.7, gel B). A quick and direct visual comparison between the lane 8 of gel A (150 $\mu$ g of TE) and the lane 1 of gel B (17.5 $\mu$ g of

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### Estimation of relative ICF-POD activity in extracts from excised gel bands.

Duplicate groups of 2 lanes of ICF-POD's 1 & 2 and 3 & 4 pairs were excised from 10% (w/v) native-PAGE gels, and treated as described in section 7.5. Guaiacol peroxidase activity was assayed in each extract. The mean total POD activity present in ICF was calculated by assaying  $35\mu g$  aliquots of freeze-dried ICF prepared in POD assay. All enzyme activities were calculated per g of fresh weight of original leaf material and the final values of extracts from bands were expressed as a percentage of the mean total POD activity in the ICF of 8.5U POD/g fresh weight (see section 4.3.4 for details of the POD assay). Data presented are the means and SD of duplicate measurements.



Assay for superoxide dismutase enzyme activity of control superoxide dismutase enzyme, TE and ICF in native-PAGE gels.

TE and ICF were prepared as described in sections 4.2.3 and 4.2.1, respectively. Both gels were 10% (w/v) native-PAGE gels and samples were prepared in native PAGE sample buffer (see section 4.4.2). The quantities quoted refer to either Bradford quantified protein concentration or the quantity of protein/enzyme activity was derived from the information of commercial suppliers (control SOD enzyme – Sigma, UK). Gel A is loaded as follows; SOD1 = 21 U (6µg) of control enzyme; SOD2 = 63 U (18µg) of control enzyme; TE = 35µg of total extract; SOD = 4.2 U (1.2µg) of control enzyme; ICF = 35µg of sodium chloride extract ICF. See section 4.4.7 for a detailed description of the SOD staining technique.



### Assay of superoxide dismutase enzyme activity in TE and ICF native-PAGE gels.

TE and ICF were prepared as described in sections 4.2.3 and 4.2.1, respectively. Both gels were 10% (w/v) native-PAGE gels and samples were prepared in native PAGE sample buffer (see section 4.4.2). The quantities quoted refer to Bradford quantified protein concentration. See section 4.4.7 for a detailed description of the SOD staining technique. Gel A is loaded left to right, as follows; Lane 1 =  $5\mu g$  TE; Lane 2 =  $20\mu g$  TE; Lane 3 =  $35\mu g$  TE; Lane 4 =  $50\mu g$  TE; Lane 5 =  $75\mu g$  TE; Lane 6 =  $100\mu g$  TE; Lane 7 =  $125\mu g$  TE; Lane 8 =  $150\mu g$  TE. Gel B is loaded left to right, as follows; Lane 1 =  $17.5\mu g$  ICF; Lane 2 =  $35\mu g$  ICF; Lane 3 =  $52.5\mu g$  ICF.

ICF) illustrated that even when 8.5 times more TE protein is loaded the distinctive bands of ICF-SOD enzyme activity observed in gel B (figure 7.7) are not present in TE. Thus, with a reasonable degree of certainty, it is highly probable that these SOD-active enzymes are apoplastic in origin.

The ICF loading study presented in figure 7.7 (gel B) clearly illustrated that when less protein is loaded it is possible to discern a distinct and reasonably well defined and very SOD active band in the upper part of the gel (best seen in lane 1), this band has been termed, ICF-SOD 4. In contrast, very distinct, well resolved and discrete bands are present in the lower half of the gel and may represent single proteins; they have been designated ICF-SOD's 1, 2, 3, and 5 (figure 7.7).

ICF's were subjected to electrophoresis through a range of acrylamide gels with decreasing pore size (figure 7.8) in an attempt to further resolve and therefore distinguish the individual ICF derived SOD-active components. It was hoped that the 5% (w/v) gel would result in better resolution of the ICF-SOD 4 band at the top of the gel. Unfortunately, it did not and the band simply broadened (figure 7.8, 5% gel). However, ICF-SOD's 1, 2, 3 and 5, were very well resolved in the 12.5% (w/v) gel (figure 7.8, 12.5% gel) and furthermore the broad region that contained ICF-SOD 4 became more discrete which made physical isolation easier for subsequent molecular weight analysis.

### 7.2.5 Spectrophotometric assay of SOD activity in excised bands

To re-confirm that ICF-SOD's 1-5 actual contained SOD activity a highly sensitive spectrophotometric enzyme assay was employed (see section 4.3.3). Briefly, ICF-SOD enzyme activity bands designated 1-5 were carefully excised from the main gel and treated as outlined in section 7.4.1, except that SOD assay buffer was used. The activity data derived from duplicate measurements were expressed as percentages of the total SOD activity present in the same quantity of protein. Similar shortcomings were evident, in that, measurement of the recovery of protein from the gel extract was not attempted, and the potential for enzyme inactivation had not been considered. Consequently, direct comparisons cannot be made, however, it appears that approximately 18% of the total SOD enzyme activity present in ICF is contained within ICF-SOD's 1-5 as separated by native-PAGE even after electrophoresis and a 24 hour incubation period (see figure 7.9).

### 7.2.6 Molecular mass estimation and sub-unit analysis of apoplastic POD's

The resolution of ICF-POD's 1, 2, 3 and 4 was deemed to be best for individual band excision and subsequent polypeptide molecular mass estimation on 10% (w/v) native



## Assay for superoxide dismutase enzyme activity in ICF separated in native-PAGE gels of differing compositions.

All gels are native (see section 4.4.2) but of differing acrylamide ratios (the actual acrylamide percentage (w/v) of each gel is indicated above the gel image). All gels were loaded with 35µg of Bradford assay quantified protein from the same sodium chloride extracted ICF (protein concentration quantified from Bradford assay results). All gels were initially stained for SOD enzyme activity; see section 4.4.6 for a detailed description of the staining technique.



### Estimation of relative ICF-SOD activity in extracts from excised gel bands.

Duplicate groups of 2 lanes of ICF-SOD1, 2, 3, 4 and 5 bands were excised from 12.5% (w/v) native gels, placed into eppendorf tubes and macerated. SOD activity was assayed in each extract. The mean total SOD activity present in ICF was calculated by assaying  $35\mu g$  aliquots of freeze-dried ICF prepared in SOD assay. All enzyme activities were calculated per g of fresh weight of original leaf material and the final values of extracts from bands were expressed as a percentage of the mean total SOD activity in the ICF of 2.2U SOD/g fresh weight (see section 4.3.5 for details of the SOD assay). Data presented are the means and SD of duplicate measurements.

gels (see figure 7.4). The method used for the transfer of POD enzyme activity stained bands from the native gel to the SDS-PAGE gel for sub-unit separation is detailed in section 7.2.

Figure 7.10 illustrates a SDS-PAGE separation of ICF-POD's 1 & 2. ICF-POD 1 appeared to be composed of 2 polypeptide components (A and B) with molecular weights of 58.1 and 29.2 kDa, which readily stained with Coomassie. Interestingly, the large polypeptide is almost exactly twice the size of the small polypeptide, this firstly indicated that the denaturation process is not 100% efficient, and secondly that ICF-POD 1 may be a dimer. ICF-POD 1 migrated to the upper portion of a native-PAGE gel (figure 7.4) and consequently it can be assumed that it either had a relatively high molecular weight and/or is of neutral pl. It is also therefore possible that component A is undenatured ICF-POD 1. If this is correct, the dimer is probably held together by disulphide linkages because it had only been partially denatured before electrophoresis. Figure 7.10 also shows the molecular weight estimation of ICF-POD 2, it is composed of one component (A) with almost exactly the same molecular weight as component B from ICF-POD1, Salzer & Hager [1993] and Otter & Polle [1997] have both described a 'cell wall' peroxidase enzyme with a molecular mass of 29 kDa from Norway spruce. It was unfortunate that neither component could be satisfactorily subjected to N-terminal sequence analysis because of N-terminal blockage.

Figure 7.11 shows the same experiment for ICF-POD's 3 and 4, no bands could be seen when the gel had been Coomassie stained. Therefore, it was counter-stained with silver stain. The higher in gel enzyme activity (figure 7.4) and spectrophotometric enzyme activity (figure 7.5) present in a relatively small quantity of material that cannot be visualised with Coomassie staining indicated that the specific activity of these polypeptides for guaiacol is much higher than that of either ICF-POD's 1 or 2 (also see figure 7.5). Both ICF-POD's 3 and 4 exhibited essentially the same polypeptide pattern, 3 components were present in both excised enzyme activity bands, it cannot be determined if these components are related or are in fact an artifact.

### 7.2.7 Molecular mass estimation and sub-unit analysis of apoplastic SOD's

12.5% (w/v) native-PAGE gels were deemed to yield the best resolution for the excision of SOD enzyme activity bands (see figure 7.4) and were used throughout this section. Figure 7.12 illustrates the initial estimation of the molecular weights of ICF-SOD's 1 and 2. ICF-SOD 1 yielded a single major polypeptide band with an apparent molecular mass of 20.2 kDa, ICF-SOD 2 generated two denatured components, A and B, with molecular masses of 25.2 and 22.2 kDa, respectively. Figure 7.12, illustrated that a low quantity of



	Component	Rf	MW (kDa)
ICF-POD1	А	0.1944	58.1
	В	0.6500	29.2
ICF-POD2	A	0.6389	29.9

### Molecular mass estimation of the sub-unit composition of ICF-POD 1 and 2.

All samples and standards were run on the same 12.5% (w/v) denaturing gel and coomassie blue stained (see sections 4.4.1 and 4.4.3, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'M' =  $6\mu$ l of Promega Mid Range molecular weight marker proteins; Lane 'ICF-SOD1' = 4x ICF-POD1 bands excised from 10% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF-POD2' = 4x ICF-POD2 bands excised from 10% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF' =  $35\mu$ g of ICF. Table was constructed from molecular weight calibration curve data (not shown). Refer to section 7.2 for details of loading strategy.



	Component	Rf	MW (kDa)
ICF-POD3	А	0.2553	55.5
	В	0.3032	52.5
	С	0.7447	25.2
ICF-POD4	A	0.2553	55.5
	В	0.3032	52.5
	С	0.7447	25.2

### Molecular mass estimation of the sub-unit composition of ICF-POD 3 and 4.

All samples and standards were run on the same 12.5% (w/v) denaturing gel, initially coomassie blue stained and the silver stained (see sections 4.4.1, 4.4.3 and 4.4.4, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'M' =  $3\mu$ l of Promega Mid Range molecular weight marker proteins; Lane 'ICF-POD3' = 1x ICF-POD3 bands excised from 10% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF-POD4' = 1x ICF-POD4 bands excised from 10% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF' = 11.7 $\mu$ g of ICF. Table was constructed from molecular weight calibration curve data (not shown). Refer to section 7.2 for details of loading strategy.


	Component	Rf	MW (kDa)
ICF-SOD1	A	0.8245	20.2
ICF-SOD2	A	0.7447	25.2
	В	0.7926	22.2

# Molecular mass estimation of the sub-unit composition of ICF-SOD 1 and 2.

All samples and standards were run on the same 12.5% (w/v) denaturing gel, initially coomassie stained and then silver stained (see sections 4.4.1, 4.4.3 and 4.4.4, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'M' =  $3\mu$ I of Promega Mid Range molecular weight marker proteins; Lane 'ICF-SOD1' = 1x ICF-SOD1 band excised from 12.5% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF-SOD2' = 1x ICF-SOD2 band excised from 12.5% (w/v) native-PAGE gel that had been activity stained; Lane 'ICF' =  $11.7\mu$ g of ICF. Table was constructed from molecular weight calibration curve data (not shown). Refer to section 7.2 for details of loading strategy.

protein was present in both ICF-SOD 1 and 2 enzyme activity stained bands because of the necessity to silver stain the SDS-PAGE gel, consequently, more native-PAGE gels were run and ICF-SOD's 1-3 re-isolated, the resultant SDS-PAGE molecular mass estimation is presented in figure 7.13. An increased loading of native-PAGE excised activity bands permitted the major polypeptides to be visualised utilising Coomassie staining only. ICF-SOD 1 and 2 were consistent and the banding patterns of polypeptides observed in figure 7.12 were successfully reproduced. A small degree of variation in the final determined molecular masses of each polypeptide was observed (~1 kDa) which presumably reflects the day-to-day variability of the system. ICF-SOD 3 produced a slightly 'wavy' single polypeptide band with an apparent molecular mass of ~22 kDa.

Scant information is available regarding plant apoplastic SOD's and the only reports that could be found detailed a Mn-SOD with a denatured molecular mass of 22 kDa from the moss, *Barbula unguiculata* [Yamahara *et al.* 1999] and a homogenous dimeric CuZn-SOD from *Pinus sylvestris* needles with a sub-unit molecular mass estimated to be around 17.8 kDa [Streller & Wingsle 1994]. Other reports of extracellular SOD's include a Fe-SOD from a cyanobacteria with a molecular mass of 21 kDa [Shirkey *et al.* 2000] and a CuZn-SOD from a nematode with a mass of 19.8 kDa [Liddell & Knox 1998]. The molecular masses reported within this thesis are of the same order as these reports even though a different species is under study thus reinforcing the native-gel and spectrophotometric assay data and confirming SOD enzyme activity.

ICF-SOD 4 was SOD-active and exhibited as a very broad band even when electrophoresed through a 12.5% (w/v) native-PAGE gel. Consequently, ICF-SOD 4 was isolated as a 3mm long piece of gel and divided into 3x 1mm pieces progressing from the top of the gel to the bottom and termed ICF-SOD4[i], [ii] and [iii], respectively. Figure 7.14 illustrates the molecular weight estimation for the polypeptides present in these gel pieces. What was immediately obvious and different from the other, highly discrete ICF-SOD's 1-3 (figure 7.13) was a diffuse low molecular weight material with apparent molecular weights of <21 kDa, that is present in all of the ICF-SOD 4 runs and that stained quite intensely. ICF-SOD 4 [i] did not exhibit a distinctive banding pattern and as such could not be analysed further, however, there was further diffuse protein material present at the top of the gel (>66 kDa) that was not present in ICF-SOD 4 [ii] or [iii] lanes. ICF-SOD's 4 [ii] and [iii] showed a doublet of bands of approximately the same mass, 44.7 kDa. The potentially multimeric, complex and diffuse nature of ICF-SOD 4 in the denaturing gel system prevented further isolation and analysis.



	Component	Rf	MW (kDa)
ICF-SOD1	A	0.8120	19.0
ICF-SOD2	А	0.6930	26.5
	В	0.7701	21.7
ICF-SOD3	А	0.7701 - 0.7594	22 ± 0.3

Molecular mass estimation of the sub-unit composition of ICF-SOD 1, 2 and 3.

All samples and standards were run on the same 12.5% (w/v) denaturing gel and coomassie stained (see sections 4.4.1 and 4.4.3, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'M' =  $6\mu$ l of Promega Mid Range molecular weight marker proteins; Lane 'ICF-SOD1' = 4x ICF-SOD1 bands excised from 12.5% native PAGE gel that had been activity stained; Lane 'ICF-SOD2' = 4x ICF-SOD3 bands excised from 12.5% native PAGE gel that had been activity stained; Lane 'ICF-SOD3' = 4x ICF-SOD3 bands excised from 12.5% native PAGE gel that had been activity stained; Lane 'ICF-SOD3' = 4x ICF-SOD3 bands excised from 12.5% native PAGE gel that had been activity stained; Lane 'ICF-SOD3' = 4x ICF-SOD3 bands excised from 12.5% native PAGE gel that had been activity stained; Lane 'ICF' =  $35\mu$ g of ICF. Table was constructed from molecular weight calibration curve data (not shown). Refer to section 7.2 for details of loading strategy.



	Component	Rf	MW (KDa)
ICF-SOD4[i]	?	Wide range	>66KDa
	?	Wide range	<21KDa
ICF-SOD4[ii]	A [doublet]	0.3249	45.9
	?	Wide range	<21KDa
ICF-SOD4[iii]	A [doublet]	0.3401	44.7
	В	0.4721	34.7
	С	0.5228	30.8
	?	Wide range	<21KDa

# Molecular mass estimation of sub-unit composition of ICF-SOD4.

All samples and standards were run on the same 12.5% (w/v) denaturing gel, initially coomassie blue stained and then silver stained (see sections 4.4.1, 4.4.3 and 4.4.4, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'M' =  $3\mu$ I of Promega Mid Range molecular weight marker proteins; Lane 'ICF-SOD4[i]' = 4x ICF-SOD4[i] bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' = 4x ICF-SOD4[ii]' bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[iii]' and the second from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[iii]' and the second from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[iii]' and the second from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF-SOD4[iii] bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF'-SOD4[iii] bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF'-SOD4[iii] bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF'-SOD4[iii] bands excised from 12.5% % native PAGE gel that had been activity stained; Lane 'ICF' = 5.85 µg of ICF. Table was constructed from molecular weight calibration curve data (not shown). Refer to section 7.2 for details of loading strategy and section 7.9 for details of how ICF-SOD4 was sectioned.

#### 7.2.8 Inhibition studies

Different types of SOD's and POD's can be broadly identified through their differential responses to inhibitors.

#### 7.2.8.1 Inhibition of ICF-SOD's

The best way to distinguish between the various types of SOD's is by the characterisation of the metal ion present [Bannister et al. 1987], cyanide is known to inhibit CuZn-SOD's but not Mn-SOD's [Fridovich 1974]; hydrogen peroxide inactivates both the CuZn- and Fe- containing SOD's but not Mn-SOD [Symonyan & Nalbandyan 1972]. Three identical loadings of ICF were run through a 12.5% (w/v) native-PAGE gel and after electrophoresis each lane was excised and treated for 5 minutes in the dark in either, 50mM phosphate buffer, pH 7.5 (control), 50mM phosphate buffer, pH 7.5 containing 1mM potassium cyanide or 50mM phosphate buffer, pH 7.5 containing 10mM hydrogen peroxide. After each treatment, the lanes were subjected to SOD enzyme activity staining. Figure 7.15 shows the result of the SOD staining, ICF-SOD's 1 and 5 were both inactivated by 1mM cyanide and by 10mM hydrogen peroxide indicating that these are probably CuZn-SOD enzymes. The activity of ICF-SOD's 2 and 3 is inhibited by treatment with hydrogen peroxide but not cyanide indicating that these are probably Fe-SOD's. Completely unaffected by either treatment was ICF-SOD 4 which was therefore assigned as a Mn-SOD. Ogawa et al. [1996] and Schinkel et al. [1998] have both isolated apoplastic CuZn-SOD's from plants, but no reports of apoplastic Fe- or Mn-SOD's could be found. However, extracellular Fe- and Mn-containing SOD's have been described in cyanobacteria [Shirkey et al. 2000] and moss [Yamahara et al. 1999], respectively.

# 7.2.8.2 Inhibition of ICF-POD's

The results of the above SOD stained inhibition study were re-confirmed in a repeat experiment using 10% (w/v) native-PAGE gels. It was decided to counter-stain these gels for ICF-POD enzyme activity to assess if any inhibition had taken place. POD's can be classified as selenium-dependent and cyanide insensitive [Paes & Oliveira 1999] or Fe-containing and cyanide sensitive [Rob *et al.* 1997]. Figure 7.16 illustrates the results of the ICF-POD inhibition study; treatment with 1mM potassium cyanide had completely inactivated all ICF-POD's, they were therefore determined to be cyanide sensitive and therefore probably were Fe-containing POD's. It is noteworthy that the TE-POD's are cyanide insensitive, marking a clear distinction between apoplastic and intracellular POD's.



# Inhibition of individual ICF-SOD enzymes.

3x lanes of  $35\mu g$  (Bradford quantified protein) loadings of the same ICF preparation were run through a 12.5% (w/v) native-PAGE gel (section 4.4.2); each lane was excised and treated individually before staining for SOD activity; buffer (control) = 50mM phosphate buffer, pH 7.5; potassium cyanide= 50mM phosphate buffer, pH 7.5 containing 1mM potassium cyanide; hydrogen peroxide = 50mM phosphate buffer, pH 7.5 containing 10mM hydrogen peroxide; each was then washed extensively for >3minutes and then SOD stained according to section 4.4.7.



# Inhibition of individual ICF-POD enzymes.

3x lanes of  $35\mu g$  (Bradford quantified protein) loadings of the same ICF preparation were run through a 10% (w/v) native-PAGE gel; each lane was excised and treated individually before staining for POD activity; buffer (control) = 5 minutes in 50mM phosphate buffer, pH 7.5 in the dark; potassium cyanide= 5 minutes in 50mM phosphate buffer, pH 7.5 containing 1mM potassium cyanide in the dark; hydrogen peroxide = 5 minutes in 50mM phosphate buffer, pH 7.5 containing 10mM hydrogen peroxide in the dark; each was then washed extensively for >3minutes and then POD stained according to section 4.4.6.

# 7.2.9 Changes in the anti-AOS status of the leaf apoplast

The activities of anti-AOS enzymes are known to alter during abiotic stress [Yahraus *et al.* 1995; Luwe 1996; O'Kane *et al.* 1996] and during development [Rogiers *et al.* 1998; Takahama *et al.* 1999]. The activities of the apoplastic forms of these enzymes were therefore assessed during abiotic stress and through development.

# 7.2.9.1 The effect of abiotic stress on ICF-SOD and POD activities

Tobacco plants were subjected to a range of abiotic conditions typically found to induce environmental stress: -

- 1 Drought was mimicked by restricting the water supply until all the leaves had wilted (~14 days) after which watering was resumed, the plants were allowed to rehydrate over a 3 day period and were then subjected to VI.
- 2 Leaves were mechanically stressed by cutting off the top fifth of one leaf from 10 tobacco plants. After 48 hours the injured leaves were collected and subjected to VI.
- 3 Temperature stress was achieved by incubating tobacco plants at 40°C in a growth chamber for 5 days, after which the leaves were subjected to VI.

Leaf ICF from each set of stressed plants was subjected to electrophoresis side-by-side and on a 10% (w/v) native-PAGE gel, and then subjected to SOD and POD enzyme activity staining, the results are displayed in figure 7.17. Although figure 7.17 provides only a visual assessment of the relative activities of each ICF-SOD and ICF-POD active band, it does illustrate that the anti-AOS enzymes present in the apoplast are differentially responsive to differing stresses.

Drought stress did not appear to have any effect on the profile or the activity of ICF-SOD's or ICF-POD's, see figure 7.17, control and drought gels. However, mechanical stress reduced enzyme activity of ICF-SOD's 1, 2 and 3 while it enhanced the enzyme activity of ICF-POD's 1 and 2. Yahraus *et al.* [1995] have observed a similar enhancement of apoplast POD activity during pathogen interaction, that may share a common signal transduction route with mechanical (wounding) stress [Jongsma *et al.* 1994; Baron & Zambryski 1995]. Elevated temperature similarly reduced the detectable enzyme activity of ICF-SOD's 1, 2 and 3, but no concurrent or dissimilar data could be



# ICF-SOD and ICF-POD patterns during abiotic stress.

All samples were run on the same 10% (w/v) native gel (see section 4.4.2) and initially stained for SOD activity and then counterstained for POD activity (see sections 4.4.7 and 4.4.6, respectively). All quantities quoted refer to protein concentrations calculated using the Bradford assay (see section 4.3.2). Lane 'Control' = 35µg of ICF VI from untreated plants; Lane 'Drought' = 35µg of ICF VI from drought stressed plants; lane 'Mechanical' = 35µg of ICF VI from mechanically stressed plants; lane 'Temperature' = 35µg of ICF VI from temperature stressed plants.

found in the literature because of the lack of studies investigating the effects of elevated temperatures.

# 7.2.9.2 The effect of leaf developmental age on ICF-SOD and POD activities

Leaves were categorised into four sets according to their relative developmental age. Care was taken to ensure that all plant material was of the same initial age by monitoring germination. 10-week old plants (post-germination) were brought together and their leaves categorised. The definitions of the different ages were as follows: -

- Age #1 defined as 'young'; excised the upper quarter of the stem.
- Age #2 defined as 'mid'; excised the next quarter of the stem.
- Age #3 defined as 'old'; excised the next quarter of the stem.
- Age #4 defined as 'senescing'; excised the bottom quarter of the stem.

Each category was excised from the stem and subjected to VI (section 4.2.2) in order to recover ICF from each leaf developmental age. ICF preparations from each leaf developmental age were prepared and loaded onto a 12.5% (w/v) native-PAGE gel, on a protein weight-for-weight basis and subjected to electrophoresis. Afterwards SOD enzyme activity staining followed by POD enzyme activity staining was undertaken, the results are presented in figures 7.18 and 7.19, gel images labelled A.

The SOD enzyme activity stained gel (figure 7.18, gel A) exhibited a general loss of SOD activity and an altered profile of ICF-SOD's as the leaves progressed through the developmental ages. This is very evident for the better separated ICF-SOD's (1, 2, 3 and 5) in the lower part of the gel, but is also evident as a general decline in ICF-SOD 4 enzyme activity in the upper part of the gel. The observable profile of ICF-SOD's using native-PAGE and in-gel enzyme activity staining, shows a rapid loss of ICF-SOD's 1 and 5 as the leaves age, until there is no detectable activity in senescing leaves (see figure 7.18, gel A, lanes 1-4). In order to quantitatively confirm the loss of ICF-SOD enzyme activity a sensitive spectrophotometric assay was utilised (section 4.3.3); the data are shown on the graph in figure 7.18 (B). The ICF-SOD enzyme activity remains stable during developmental ages 1 and 2, but rapidly drops to less than 40% of this value by the time the leaves are undergoing senescence.

Similarly, to detect if ICF-POD enzyme activities had altered the native-PAGE gel was counter stained for POD enzyme activity, see figure 7.19 (A). The gel image shows that during early development (lane 1) and into maturity (lane 3) there was very little discernible difference between the ICF-POD enzyme profile and apparent activity using



# Quantitative analysis of ICF-SOD enzyme activity during leaf development.

A = All samples were run on the same 12.5% (w/v) native gel (see section 4.4.2) and initially stained for SOD activity (see section 4.4.7). All lanes were loaded with  $35\mu g$  of ICF protein as calculated from Bradford assay results (section 4.3.2). Refer to section 7.12 for details of how leaf age was determined. ICF was prepared from leaves of differing developmental stages as described in section 4.2.1.

B = Refer to section 7.12 for details of how leaf age was determined. ICF was prepared from leaves of differing developmental stages as described in section 4.2.1. Duplicate lyophilised solid equivalent to  $35\mu g$  of protein from each developmental stage was reconstituted in SOD assay buffer (50mM phosphate, pH 8.2) and assayed as per section 4.3.5. Units of activity were scaled back to reflect the quantity of tissue the ICF had been sampled from.



# Quantitative analysis of ICF-POD enzyme activity during leaf development.

A = All samples were run on the same 12.5% (w/v) native gel (see section 4.4.2) and initially stained for POD activity (see section 4.4.6). All lanes were loaded with  $35\mu g$  of ICF protein as calculated from Bradford assay results (section 4.3.2). Refer to section 7.12 for details of how leaf age was determined. ICF was prepared from leaves of differing developmental stages as described in section 4.2.1.

B = Refer to section 7.12 for details of how leaf age was determined. ICF was prepared from leaves of differing developmental stages as described in section 4.2.1. Duplicate lyophilised solid equivalent to  $35\mu g$  of protein from each developmental stage was reconstituted in POD assay buffer (100mM phosphate, pH 7.0) and assayed as per section 4.3.4. Units of activity were scaled back to reflect the quantity of tissue the ICF had been sampled from.

native-PAGE and in gel enzyme activity staining. However, contradictory data are displayed on the graph (figure 7.19, B), that shows a constant decrease in total ICF-POD enzyme activity as the leaf tissue progresses through each of the developmental ages. It is therefore possible that the gel (A) had been over-stained for POD activity. What is obvious from both the gel image (A) and the spectrophotometric assay (B) is that total ICF-POD activity is greatly reduced when the leaves are undergoing senescence.

A similar age-related decrease in enzymatic anti-AOS capacity has been observed previously [Rogiers *et al.* 1998; Takahama *et al.* 1999] and will be expanded upon in the conclusions (section 7.9).

# 7.2.9.3 The effect of leaf developmental age on the level of apoplastic oxidative protein modification.

In the introduction to this thesis (section 1.5) it had been outlined that the apoplast has a number of important anti-AOS strategies, broadly they can be classified into one of two depending upon their mode of action, i/ AOS-scavengers, or, ii/ anti-AOS enzymes. (see table 1.5). The above data had revealed that the activities of the anti-AOS enzymes ICF-SOD's and ICF-POD's declined as leaves aged, consequently it was inferred that the level of AOS present would gradually increase in an age-dependent manner.

In order to test this assumption the degree of oxidative modification to proteins and polypeptides (one of the major targets of AOS) can be assessed using an antibody based assay specific for the oxidative modification of proteins induced by the oxidative action of the hydroxyl radical. This is the 'oxyblot kit' described in section 4.4.10.

Equal quantities of ICF Bradford assay quantified (section 4.3.2) protein from the differing leaf developmental ages were therefore analysed for oxidative protein modification and the data are presented in figure 7.20. A general increase in the level of oxidative protein modification, which is specific for AOS, can be seen in the apoplastic extracts as the leaf material progresses through the developmental ages. These data follow the trend of decreasing ICF-SOD and ICF-POD activities observed in figures 7.18 and 7.19 and it is thought that the decreased capacity to detoxify AOS directly elevates the number of oxidative protein modifications.



# Oxidative modification of ICF proteins during leaf development.

ICF's were prepared from leaves at different stages of development as described in section 7.12. Samples and standards were derivatised and run through a 10% (w/v) SDS-PAGE gel (section 4.4.1), blotted to nitrocellulose membrane (section 4.4.9) and antibody probed for oxidative modification to proteins using a commercial kit (Oncor, France; see section 4.4.10); a horseradish peroxidase conjugated secondary antibody was used for detection. Samples are labeled on the oxyblot as progressing from the youngest leaves (#1) through to the senescing (#4). All leaf ICF samples were at 11.7 $\mu$ g of protein and the remainder of the lanes were loaded as follows; Lane 'TE' = 11.7 $\mu$ g; Lane 'OxyM' = 5 $\mu$ l of oxidatively modified marker proteins. All quoted sample concentrations have been

#### 7.4 Conclusions

Utilising native-PAGE, in gel enzyme activity assays and more specific spectrophotometric enzyme assays this part of the study has identified that the tobacco leaf apoplast contains POD's and SOD's that cannot be found inside the cell.

Four distinct guaiacol POD utilising enzyme active bands, which display unique separation patterns in native gels indicating that they have differing properties were visible and clearly not present in TE preparations. Molecular mass estimation revealed that ICF-POD 1 was potentially a disulphide linked dimer composed of two sub-units with identical molecular mass of ~29 kDa, both Salzer & Hager [1993] and Otter & Polle [1997] have reported the isolation of a basic cell wall 'bound' POD enzyme with the same molecular mass as the sub-unit of ICF-POD 1. All of the four identified ICF-POD's were designated as Fe-containing POD's rather than selenium dependent because of their sensitivity to the cyanide ion [Rob et al. 1997] and the activity of each was observed to reduce in an age-dependent manner. Very interestingly ICF-POD's 1 & 2 co-migrated in the native-PAGE system with a broad SOD-active band, ICF-SOD 4, and although in an entirely unrelated system, a recent report from Johansson et al. [1999] details a cell surface POD that can ionically interact with an extracellular SOD that can only be extracted from cells when treated with a salt solution. It is postulated that the POD-SOD interaction will enhance the efficiency of AOS breakdown via closer proximity of the POD to its substrate. A similar arrangement of anti-AOS enzymes at the plasma-membrane would be beneficial in preventing inadvertent membrane damage from apoplastic AOS.

Five distinct apoplastic SOD active bands were identified, which display unique separation patterns in native gels indicating that they have differing properties. As stated in section 7.2, it is impossible to assign distinct physical characteristics to proteins in native separation systems, but generalisations can be made. ICF-SODs 1, 2, 3 all migrated far into the gel indicating that they are probably small proteins with relatively high pl's. ICF-SOD's 1, 2 and 3 were indeed shown to be composed of small polypeptides ranging between 19 and 25 kDa. Furthermore, ICF-SOD 1 was designated as a CuZn-SOD based on its sensitivity to both cyanide and hydrogen peroxide, Streller *et al.* [1994] and Schinkel *et al.* [1998] have similarly identified apoplastic SOD's from Scots pine needles that have approximately the same molecular mass, are CuZn-SOD's and that have high isoelectric points, it seems likely that ICF-SOD 1 is a very similar tobacco enzyme. The inhibition study also indicated that ICF-SOD 4 is probably a Mn-SOD and that ICF-SOD's 2 and 3 are probably are Fe-containing SOD's. The first plant extracellular Mn-SOD was only identified within the last year [Yamahara *et al.* 1999] and to the authors knowledge the presumed Fe-containing apoplastic SOD's (ICF-SOD's 2 &

3) are the first to be described in plants. An extracellular Fe-SOD has very recently been discovered in bacteria [Shirkey *et al.* 2000] but not in higher organisms and interestingly, it has almost the same molecular mass as ICF-SOD 3. The overall ICF-SOD activity was observed to reduce in an age-dependent manner and the rapid and specific loss of the CuZn-SOD's (ICF-SOD's 1 and 5) was very evident.

The study presented here shows a distinct decline in the activity of SOD and POD extracted from the apoplast, thus indicating that the overall antioxidant capacity of the apoplast also declines in a similar manner to that of the intracellular environment during aging. It was proposed a number of years ago [Thompson 1984] and more recently established [Ferrie et al. 1994; Rogiers et al. 1998] that the maturation and ripening of fruit is accompanied by a general decline in the ability to cope with AOS. A very recent, and more relevant study by Takahama et al. [1999] has detailed a general decrease in leaf tobacco apoplastic anti-AOS compounds, such as ascorbate, depending upon the position of the leaf on the plant (i.e.: the age of the leaf). Surprisingly though, some of the ICF-SOD and ICF-POD activities measured by Takahama et al. [1999] predominantly increased as the leaves aged, and then dropped rapidly during senescence. The discrepancy between Takahama et al. [1999] and the results displayed herein may be artifactual and arise because of the different methods used for grouping leaves into a particular developmental age. An oxyblot analysis confirmed that an age-dependent loss of apoplastic anti-AOS enzymes results in an age-dependent increase in the levels of hydroxyl radical mediated protein modification, similar results have been obtained for lipid peroxidation [Thompson 1984].

#### **SECTION 8**

#### RADICAL RESPONSES TO STRESS IN THE APOPLAST

#### 8.1 Introduction

The production of active oxygen species (AOS) in the apoplast during plant-pathogen interactions has been well documented [see Mehdy 1994; Wojtaszek 1997 for reviews, section 1.5 and figure 1.5]. In addition to biotic stress, chemical [Kangasjarvl *et al.* 1994], temperature [Fadzillah *et al.* 1996; O'Kane *et al.* 1996], osmotic [Cazalé *et al.* 1998] and mechanical stresses [Jaffe *et al.* 1985; Legendre *et al.* 1993; Yahraus *et al.* 1995] can all induce the production of AOS (see section 1.5 and figure 1.6) although the mode of production is still under investigation. Typically, these studies have quantified the production of hydrogen peroxide from cell suspension cultures as an indicator of the presence of the other AOS. Unfortunately, the use of cell suspension cultures reduces the physiological relevance of the data and the measurement of hydrogen peroxide rather than the 'precursor' molecule, the superoxide anion (O<sub>2</sub>), is not stress-specific (see figure 1.6). The direct quantification of O<sub>2</sub><sup>-</sup> in leaves during exposure to ozone has been documented [Runeckles & Vaartnou 1997] but required the use of an expensive electron paramagnetic resonance spectrometer and highly specialised knowledge for the interpretation of the resultant data.

Nitric oxide (NO) is an integral component of the inflammatory signalling pathway leading to apoptosis in animals [Kooy *et al.* 1995] and it has recently been postulated that the cell death observed during the hypersensitive response of plants to pathogens is essentially the same mechanism [see Douglas 2000, for an overview]. The first report of nitric oxide involvement in plant defence responses came after Noritake *et al.* [1996] illustrated that NO can induce phytoalexin accumulation in potato. In 1998 Delledonne *et al.* and Durner *et al.* both independently established that the production and release of NO and AOS was integral to hypersensitive cell death during plant-pathogen interactions. Interestingly it is the interplay between NO and  $O_2^-$  to generate the peroxynitrite radical that is proposed to mediate programmed cell death [Sadoval *et al.* 1997], exert direct anti-microbial effects [Mayer & Hemmens 1997] and modulate the activity of a range of other proteins [Ischiropoulos *et al.* 1992; Jiamrong *et al.* 1997; So *et al.* 1998; Souza & Radi 1998]. A more detailed description is provided in section 1.5.4.

The apoplast is central to the interaction of NO and AOS due to the fact that it is one of the primary sites for the generation of AOS [Mehdy 1994; Wojtaszek 1997; Scheel 1998]

and because many of the potential effects of NO and the resultant peroxynitrite radical are postulated to occur in the extracellular environment [Delledonne *et al.* 1998; Durner *et al.* 1998; see section 1.5]. As a consequence, investigation of the apoplastic presence of NO and  $O_2^-$  during stress was undertaken.

Throughout this project VI had been utilised as an effective method for selectively sampling proteins and polypeptides from the leaf apoplastic environment. However, the technique was not applied to the study of apoplastic NO and the superoxide anion primarily because apoplastic fluid from tobacco leaf tissue typically had a pink colouration that would have directly interfered with the chosen assay systems and secondly because the lag period between infiltration and recovery was deemed to be too variable. A series of experiments utilising oxyblot analysis (section 4.4.10) had been carried out by S. Johnson [Croy *et al.* 1998; and personal communications with S. Johnson] at the University of Durham. These experiments had inferred that impact stressed potato tuber tissue generated AOS from a small, well defined sub-epidermal site directly below the point of impact. The tuber-impact experimental system offered several important advantages for the study of apoplastic  $O_2^-$  and NO in comparison with the standard leaf VI model:-

- 1. It was important to know that this system had the potential to yield positive results; Croy *et al.* [1998] had already indirectly shown that AOS were present.
- Because AOS generation in response to impact stress was limited to a very small site it is technically easier to measure AOS/NO generation furthermore, it was possible to excise control tissue from the same tuber and thus reduce any potential variability that may be present between tubers.
- 3. Providing that the AOS generating site was carefully excised and extensively washed (in a similar manner to leaf pieces during the VI method), it could be assumed with a reasonable degree of certainty that AOS/NO were being produced and/or released in the apoplast in the intact tuber.
- 4. The reagent used for the quantification of the superoxide anion is expensive; using the tuber-impact system would greatly reduce the costs because relatively small volumes of reagent would be needed.

Consequently, the tuber-impact model system was utilised during this research for the study of apoplastic  $O_2^-$  and NO.

The initial work describing the design and optimisation of assays for the detection of the superoxide anion and nitric oxide were originally designed for use with mechanically stressed cell suspension cultures and were carried out solely by the author. Subsequently the assays were applied to a more biologically relavent system, namely the response of potato tubers to impact stress. This latter experimental project was jointly developed with PhD student colleague (Mr. S. Johnson). The effort in this part of the work was equal and has subsequently been submitted as a joint publication to Planta (see appendix 1).

#### 8.1.2 Notes on experimental design

An impact stress of 0.7 joules was applied to one end of the tuber by dropping a bolt of defined weight from a pre-determined standard height. The impact time was designated as time zero and the impact site was marked with a felt-tipped pen. The tuber was either used immediately (zero time point) or incubated for a required period. After incubation the tuber was halved through the impact site with a sharp knife and duplicate half cores (9mm long by 2.5mm radius) were taken from each half of the tuber. Duplicate control cores were excised at the 2 and 10 o'clock positions with respect to the impact site cores (see figure 8.1). Each core was cut into 3x 3mm long sections with a sharp blade and washed extensively for >5 minutes with at least 3 changes of dH<sub>2</sub>O to remove any leaked intracellular components. See figure 8.1 for a diagrammatic overview. Tuber pieces were then thoroughly dried with paper towel, placed into individual eppendorf tubes and submerged under  $200\mu$ l of the radical detection reagents. The tubes were placed in the dark for the 20 minute assay period since both of the radical detection systems have light sensitive components. After 20 minutes the tuber pieces were removed and discarded, and the assay reagents were centrifuged for 5 minutes in a benchtop microfuge (13000g) and the optical density of the supernatants measured against the appropriate reagent blank (see sections 8.2.1 and 8.2.2 for details of each assay system).

#### 8.2 Results

#### 8.2.1 Assay development

In order to quantify the generation of the superoxide anion and nitric oxide from potato tuber tissue assays for the quantitative detection of each molecule were developed.

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Experimental Outline : Impact stress and sample collection. Image compiled for this thesis; see section 8.1.2 for a detailed explanation.

# 8.2.1.1 Development of an assay system for the detection of the superoxide anion (O<sub>2</sub>)

The superoxide anion can be measured directly because of its distinctive physical characteristics; an unpaired electron generates a diagnostic electron paramagnetic resonance spectrum, and unique UV absorption [Marklund 1976]. Unfortunately, neither of these methods is suitable for routine quantification of the superoxide anion in biological systems. The redox protein, cytochrome C has been utilised for superoxide quantification previously, but was not chosen for this study because it has a molecular weight of 12.5 kDa which would hinder efficient diffusion through cell walls to the proposed site of superoxide production, the outer face of plasma-membrane. The hydroperoxyl/superoxide anion acid-base pair ( $O_2H/O_2^-$  - see section 1.5.1 for details of the chemistry of AOS) selectively reduce tetrazolium compounds producing formazan dyes that can be readily quantified.

#### Tetrazolium Salt + $O_2^*$ ----- Formazan dye ( $\lambda_{max}$ )

Nitroblue tetrazolium has been utilised for the colourimetric detection of the  $O_2H/O_2$  pair, unfortunately, the resulting formazan is relatively insoluble in aqueous solutions, a highly desirable quality for histochemical localisation studies, but less suitable for kinetic or simple quantification studies. Relatively recently a new tetrazolium compound, XTT (2,3-bis-[2-Methoxy-4-nitro-5-sulphophenyl]-2H-tetrazolium-5-carboxanilide) had been utilised successfully for the measurement of  $O_2H/O_2^-$  in cell suspension cultures during the pathogen-induced oxidative burst [Sutherland *et al.* 1998]. The reduced form of XTT is fully soluble in aqueous solutions thereby obviating the insolubility problems associated with nitroblue tetrazolium, it is also relatively small compared with cytochrome C and consequently will not encounter any diffusion barriers.

The vast majority of AOS-measurement studies have been limited to the quantification of hydrogen peroxide as an indicator of the  $O_2H/O_2^-$  radical pair, even though hydrogen peroxide is a relatively ubiquitous substrate and by-product of many intracellular and apoplastic processes [Willekens *et al.* 1995; Schopfer 1996]. XTT has been proposed to react selectively with the  $O_2H/O_2^-$  radical pair [Ewing & Janero 1997] therefore any inadvertent  $H_2O_2$ -mediated reduction of XTT (i.e.: cross-reactivity) would elevate the estimated concentration of  $O_2H/O_2^-$ , effectively increasing the level of false positive error in any measurements. Consequently, the cross reactivity of  $H_2O_2$  was assessed by spectrophotometrically following XTT reduction in the presence of  $H_2O_2$  only. Figure 8.2 depicts the data; as can be seen, there was no significant change in optical density at



# Hydrogen peroxide reaction with XTT.

 $200\mu$ I of 50mM phosphate buffer, pH 7.8 containing,  $100\mu$ M XTT was pipetted into the wells of a microtitre plate. Stock hydrogen peroxide was diluted to form a calibration series of hydrogen peroxide concentration at 0.2, 0.5, 1, 2 and 20mM/mI; a 50 $\mu$ I aliquot of each was added to the XTT solution at time zero and the microtitre plate was then stored in a light proof container until OD<sub>450</sub> (optical density) was measured after 20 minutes against an XTT control lacking H<sub>2</sub>O<sub>2</sub>. The final concentration of hydrogen peroxide per mixture is show on the abiscca. Data are the means of ± SD of four measurements.

 $\lambda_{max}$  for the reduced formazan over a 20-minute period, thereby indicating that hydrogen peroxide and XTT do not cross-react.

In order to verify the efficacy of the O<sub>2</sub>H/O<sub>2</sub>-induced reduction of XTT, a method to artificially generate O<sub>2</sub> radicals was employed. Commonly, superoxide generating systems utilise the enzyme xanthine oxidase however, xanthine oxidase alone can also induce reduction of related tetrazolium compounds (e.g.: nitroblue tetrazolium) [Beauchamp & Fridovich 1971] thus causing an increase in false positive error, consequently it was not chosen for this study. Instead a non-enzymatic system of superoxide anion production was adopted utilising aerobic mixtures of NADH and phenazine methosulphate (PMS) [Ewing & Janero 1997].

[red]PMS + O<sub>2</sub> + NADH  $\longrightarrow$  [ox]PMS + O<sub>2</sub> + NAD<sup>+</sup> + H<sup>+</sup>

It is known that the superoxide anion generated from either biological material or chemically via the PMS-NADH system dismutates to hydrogen peroxide and oxygen with increasing rapidity from basic through to acidic pH [Oberly & Spitz 1996]. In order to ensure accurate assay of the superoxide generated from either biological material or chemically via the PMS-NADH system, the pH was adjusted to ensure the longevity of the O<sub>2</sub>H/O<sub>2</sub> pair and therefore to maximise the chance of XTT reduction. Figure 8.3 depicts the results of this experiment. At the pH extremes measured (<7.5 and >8.7), XTT reduction dropped markedly indicating that the O2H/O2 pair were dismutating at a faster rate or that the PMS-NADH system was not producing as much superoxide at these pHs. XTT reduction was maximal between pH 7.5 and 8.7 indicating that there was a stable concentration of the  $O_2H/O_2$  pair in the reaction mixture and that the reduction of XTT was occurring at a relatively constant rate. A proportion of the superoxide may be dismutating into hydrogen peroxide and oxygen, but a 'fixed' quantity is reacting with XTT. The remainder of the experiments for setting up the superoxide detection assay and those described later with sample material were carried out at pH 8.2 to ensure maximal reaction rate between superoxide and XTT.

When attempting to quantify the  $O_2H/O_2$  from intact biological samples it was essential to ensure that the concentration of XTT (i.e.: the radical scavenger) was saturated. In this state the reduction of XTT and hence colour formation by the superoxide anion is maximal. Reacting varying concentrations of the radical scavenger, XTT with the standard PMS-NADH-superoxide generating system assessed this, figure 8.4 shows the data. The reaction rate between superoxide and XTT, as determined by optical density change per minute, was shown to be greatest at XTT concentrations of  $\geq 120\mu$ M, illustrating that the reaction was proceeding at a maximal rate. XTT in the PMS-NADH-



# The reaction between XTT and superoxide with varying pH.

200 $\mu$ l of 50mM phosphate buffer, pH in the range 6.5-8.9, containing, 0.1mM EDTA, 100 $\mu$ M XTT and 98 $\mu$ M NADH were pipetted into the wells of a microtitre plate. Superoxide radical production was initiated with 50 $\mu$ l of 14 $\mu$ M PMS in 50mM phosphate buffer, pH in the range 6.5-8.9. OD<sub>450</sub> was measured every 30 seconds over 3 minutes from which  $\Delta$ OD<sub>450</sub>/minute was calculated. Then data are the means of ± SD of four measurements.



# Determination of optimal XTT concentration for superoxide detection.

50mM phosphate buffer, pH 8.2 containing,  $80\mu$ M EDTA,  $78.4\mu$ M NADH and  $6-150\mu$ M XTT. The reaction was initiated with a final PMS concentration of  $2.8\mu$ M. The optical density was read at 2 minutes from which  $\Delta$ OD<sub>450</sub>/minute was calculated. The final concentration of XTT/assay is displayed on the abscissa. Data are means of ± SD of four measurements.



# Progress of radical generation by the PMS-NADH system.

 $200\mu$ l of 50mM phosphate buffer, pH 8.2 containing, 0.1mM EDTA, 150 $\mu$ M XTT and 98 $\mu$ M NADH were pipetted into the wells of a microtitre plate. Superoxide radical production was initiated with 50 $\mu$ l of varying concentrations of PMS in 50mM phosphate buffer, pH 8.2. OD<sub>450</sub> was measured every 30 seconds over 4.5 minutes. The key for the symbols, the final concentration of PMS/assay and the resultant correlation coefficients ( $r^2$ ) for 180 and 275 second measurements is displayed in the box on the graph. The data are the means of ± SD of four measurements

superoxide generating system at concentrations  $\ge 120\mu$ M is effectively saturated with respect to its reaction with superoxide and consequently it is acting as an efficient radical scavenger.

Preliminary investigations using mechanically stressed tuber tissue indicated that in order to obtain a reasonably measurable optical density change (set at ~0.1 △OD U) tuber tissue needed to be incubated for approximately 20 minutes (data not shown). During this assay period it was necessary to ensure that the concentration of XTT did not fall below the saturation concentration of  $120\mu M$ . Figure 8.5 illustrates the concentration-dependent superoxide production by the PMS-NADH system that is ultimately dependent upon the electron donor, PMS. At a PMS concentration of 2.8µM linearity of XTT reduction is evident for the first 3 minutes of the reaction ( $r^2$ =0.9920, p<0.0001), illustrating that during this period there is enough XTT to react with the radicals at a maximal rate. Beyond 3 minutes linearity is lost ( $r^2$ =0.9613, p<0.0001) indicating that either the generation of the superoxide anion has decreased or that the XTT concentration has dropped below the saturation level of  $120\mu M$ . Adding additional XTT to the reaction mixture results in a resumption of optical density increase showing that it is not superoxide generation that has slowed but XTT reduction (data not shown). Using the published extinction coefficient for XTT at  $\lambda_{max}$  of 2.16 x 10<sup>4</sup> M<sup>-1</sup> cm<sup>-1</sup> [Sutherland & Learmonth 1997] and the change in optical density reading of 0.125  $\Delta_{450}$ /min it can be calculated that 4.34 nMoles of  $O_2H/O_2^$ reacted with XTT before its concentration drops below the saturation threshold of 120µM. Providing the biological system under study does not produce more than 4.34 nMoles in the 20 minute assay period an XTT saturation state will be maintained. Able et al. [1998] quantified O2H/O2 generation from tobacco cell suspension cultures using XTT, and calculated that 0.6 nMoles of O<sub>2</sub>H/O<sub>2</sub>/g fresh weight/min, scaling to 20 minutes gives, 12 nMoles  $O_2H/O_2/g$  fresh weight. This is greatly exceeds the saturation range of XTT calculated within this work, but it should be noted that the tissue pieces used within these experiments were much smaller than one gram and weighed of the order 30-35mg. Consequently reducing the XTT concentration below the saturation level should not occur providing a 35mg piece of tuber tissue does not generate >4.34 nMoles  $O_2H/O_2^{-1}$  over a 20 minute assay period, this effectively equates to ~124 nMoles O<sub>2</sub>H/O<sub>2</sub>/g/20 minutes, a very large quantity of superoxide ! The final assay set-up is concisely described in section 4.3.5.

# 8.2.1.2 Development of an assay system for the detection of the nitric oxide (NO)

A number of colourimetric, fluorometric and chemiluminescent assays for quantification of nitric oxide (NO) released from S-nitrothiol compounds (see section 1.5.4) have been

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Figure 8.6The reaction of the Neutral Griess reagent with 'NO.Adapted from Cook et al. 1996.

described previously [Cook *et al.* 1996]. The method chosen relies upon the spontaneous reaction of NO and molecular oxygen to form nitrite compounds (NO<sub>x</sub>) which react with the neutral Griess reagent (composed of sulphanilamide (SULF) and N-(1-napthyl)ethylenediamine dihydrochloride (NEDD)) to form an azo dye [Nims *et al.* 1995]. A simplified reaction scheme is presented in figure 8.6. Utilising this reaction has several distinct advantages over other methods of quantification, importantly no extremely toxic or hazardous chemicals are used, the reaction occurs maximally at neutral pH ( $\pm$ 1 pH unit) and finally, no major alterations to the experimental set-up described in section 8.1.2 are required. Consequently, because measurements for both NO and the O<sub>2</sub> are taken in exactly the same manner they will be directly comparable and error introduced by altering the experimental system to accommodate a different type of measurement will be greatly reduced.

A major consideration when considering the independent measurements of the superoxide anion and nitric oxide was whether the independent assay systems would cross-react. Figure 8.7 confirms that superoxide anions generated by the PMS-NADH system do not cross-react with the SULF-NEDD NO detection system thereby illustrating that azo dye formation and consequent colour change will reflect NO concentration only.

In order to verify the efficacy of NO detection using the SULF-NEDD assay a chemical means of NO production was employed. Sodium nitroprusside (SNP) releases NO when exposed to a reducing agent such as dithiolthreitol (DTT) [Delledonne et al. 1998]. Figure 8.8 depicts the production profiles of NO from varying concentrations of SNP by 10mM DTT utilising the SULF-NEDD assay system over a 20-minute period. After an 18-minute period absorbance plateaued at all SNP concentrations, effectively all the NO had been released and had reacted with the SULF-NEDD reagent. This can be illustrated by the addition of more SNP - the optical density increase is effectively resumed (data not shown). Taking these data and plotting them as a 20-minute end point optical density at  $\lambda_{max}$  (figure 8.9) it is immediately obvious that the reaction is linear, with respect to NO chemically released from SNP ( $r^2=0.9975$ ; p<0.001), the reaction is therefore maximal and the assay is quantitative. Using the published extinction coefficient for the SULF-NEDD reagent of 12,500 M<sup>-1</sup> cm<sup>-1</sup> [Nims et al. 1995] the 400µM and 100µM SNP concentrations are releasing approximately 3.5 nMoles and 0.4 nMoles of NO over the 20 minute period respectively, these figures are in approximate concordance with published concentrations of NO produced by plant systems [Delledonne et al. 1998]. The final assay set-up is concisely described in section 4.3.6.



# Superoxide anion cross-reaction with the SULF-NEDD reagent.

100 $\mu$ l of 50mM phosphate buffer, pH 7.8 containing 114mM SULF and 2.4mM NEDD were pipetted into the wells of a microtitre plate, to each a further 100 $\mu$ l of 50mM phosphate buffer, pH 7.8 containing 0.2 $\mu$ M EDTA and 196 $\mu$ M NADH were added. At time zero a 50 $\mu$ l aliquot of varying concentrations of PMS dissolved in 50mM phosphate buffer, pH 7.8 was added to each well in order to initiate superoxide anion generation. The microtitre plate was then stored in a light proof container until OD<sub>496</sub> was measured at time 20 minutes against a blank in which PMS was replaced was buffer. The final concentration of PMS per mixture is show on the abiscca. The data are the means of  $\pm$  SD of four measurements





#### Azo dye formation produced by 'NO liberated from SNP.

100 $\mu$ I of 50mM phosphate buffer, pH 7.8 containing 114mM SULF and 2.4mM NEDD were pipetted into the wells of a microtitre plate, to each a further 100 $\mu$ I of 50mM phosphate buffer, pH 7.8 containing varying concentrations of SNP was added and mixed. At time zero a 50 $\mu$ I aliquot of 50mM phosphate buffer, pH 7.8 containing 50mM DTT was added to each well in order to initiate NO liberation from SNP. OD readings at 490nm were taken every 2 minutes over a 20 minute period versus a buffer control (lacking DTT), between readings the microtitre plate was stored in a light proof container. The final concentration of SNP in each mixture is shown in the box. The data are the means of ± SD of four independent measurements.



Detection of reaction between SULF-NEDD and NO liberated from SNP at different concentrations.

100 $\mu$ l of 50mM phosphate buffer, pH 7.8 containing 114mM SULF and 2.4mM NEDD were pipetted into the wells of a microtitre plate, to each a further 100 $\mu$ l of 50mM phosphate buffer, pH 7.8 containing varying concentrations of SNP was added and mixed. At time zero a 50 $\mu$ l aliquot of 50mM phosphate buffer, pH 7.8 containing 50mM DTT was added to each well in order to initiate NO liberation from SNP. OD was measured at 490nm after 20 minutes versus a control (lacking DTT); the microtitre plate was stored in a light proof container until read. The final concentration of SNP in each mixture is shown in the box. The data between 100 and 400 $\mu$ m of SNP have been subjected to linear regression, the data is presented in the box on the graph. The data are the means of ± SD of four independent measurements.



# Measurement of the superoxide anion produced by impact stressed tuber tissue.

Experimental design described in section 8.1.2 and the final assay is presented in section 4.3.5. Briefly, the tuber was subjected to impact, the impact site was then marked and the potato incubated for 4 hours at room temperature. Duplicates cores were taken from both the impact site and a control site and incubated in the 50mM phosphate buffer, pH 8.2,  $150\mu$ M XTT; in the presence or absence of 100U of SOD (Sigma, UK), at ambient temperature in the dark. Tuber tissue was removed from the reagents after 20 minutes and discarded, the reagents were then centrifuged for 5 minutes and the OD of the supernatant at 450nm measured. The data are the means of  $\pm$  SD of 4 measurements. The inset graph illustrates the decreasing linear response of superoxide production from tuber tissue as cores are taken deeper into the impact stressed tuber; the data was derived from 'Impact site: No SOD' data presented on the main graph. The key (top right) indicates depth of core into the tuber.

#### 8.2.2 Superoxide anion quantification from tuber tissue

Mechanical stress has been stated as a trigger for the generation of AOS in plants [Jaffe *et al.* 1985; Legendre *et al.* 1993; Yahraus *et al.* 1995] and specifically potato [Croy *et al.* 1998]. A preliminary study was carried out to test the superoxide assay developed in this study using mechanically stressed tuber tissue (c.v. King Edward), figure 8.10 depicts the data. Superoxide anion production was shown to be approximately four times as great at the impact site at 4 hours post impact stress when compared with the control site. This can be reduced to level of the control site by the addition of 100U of the anti-AOS enzyme superoxide dismutase (see section 1.5.3). This therefore confirms that it is the superoxide anion that is causing the optical density change. The control site manifests approximately 25% of the superoxide generation that the impact site exhibits. It is assumed that this high background level of superoxide anion generation is induced by the general trauma and mechanical stress imposed upon the tuber in order to recover the core for quantitative superoxide measurement. This background level of superoxide anion generation with 100U of superoxide anion generation can effectively be reduced to zero by incubation with 100U of superoxide dismutase.

These data effectively illustrate that we have a largely localised and specific generation of the superoxide anion at the impact site that is preferentially confined to the outer 3mm of the tuber (see figure 8.10). Progressing a few millimetres more into the tuber and the production of superoxide tails off indicating a spatial regulation of superoxide generation. The inset graph on figure 8.10 illustrates that this constant reduction of superoxide generation bears a linear relationship with depth into the tuber core ( $r^2$ =0.9902; p<0.1). Consequently, in order to maximise and standardise all subsequent superoxide measurements they were conducted using only the outer 3mm of tuber tissue.

Figure 8.11 depicts the data derived from a preliminary time course experiment using c.v. King Edward potatoes and shows that between 4 and 6 hours the impact site generates upto 4 times more superoxide than the control site. During this period the difference in superoxide generation between the impact and control sites is statistically significant (p<0.05) and clearly illustrates that the generation of the superoxide anion is regulated with respect time.

Even though subsequent repeats of this experiment re-confirmed the general trend of greater superoxide anion generation from the impact site (data not shown), the data were generally more variable. It is well documented that the nutrient supply during growth, tuber age, storage temperature [see Peters 1996 for a review] and previous impact status [S. Johnson, personal communication] all have a profound effect upon the ability of tuber material to respond to impact stress. The tuber material utilised for these trial experiments



t=1.721 df=5

ം

Number of pairs

t, df

Figure 8.11

Preliminary time course of superoxide anion production from impact stressed tuber tissue.

Experimental design described in section 8.1.2 and the final assay is presented in section 4.3.5. The experiment was carried out using shop bought King Edward potatoes and the data is presented on the graph. The data from the control site and the impact site were statistically assessed (using GraphPad Prism software) to evaluate if there was any significant difference between the two data sets; this is shown below the graph. P-value summary key; ns=no significant difference; \*=significant difference; \*=very significant difference.



# Nitric oxide production from impact stressed tuber tissue.

Experimental design described in section 8.1.2 was used and the final assay is presented in section 4.3.6. Briefly, the tuber was subjected to impact, the impact site was then marked and the potato incubated for 4 hours at room temperature. Duplicates cores were taken from both the bruise site and a control site and incubated in the 50mM phosphate buffer, pH 7.8, containing 114mM SULF and 2.4mM NEDD in the dark. Tuber tissue was removed from the reagents after 20 minutes and discarded, the reagents were then centrifuged for 5 minutes and the OD of the supernatant at 496nm measured. The data are the means of  $\pm$  SD of 4 measurements. The inset graph illustrates the decreasing production of nitric oxide production from tuber tissue as cores are taken deeper into the impact stressed tuber; the data was derived from 'Impact site: No SOD' data presented on the main graph. The key (top right) indicates depth of core into the tuber.


had been bought from a local supermarket outlet and consequently were not of known history.

# 8.2.3 Nitric oxide quantification from tuber tissue

A preliminary experiment to establish if NO generation and consequent quantification could be achieved in intact tuber tissue (c.v. King Edward) and to indicate the degree and type of spatial regulation of nitric oxide generation was undertaken in the same manner as for superoxide anion detection (section 8.2.2). Figure 8.12 depicts the data. In a broadly similar way to superoxide anion generation observed in figure 8.10, NO appears to be spatially regulated, with higher quantities found locally at the impact site when compared with the control site. Furthermore, the quantity of NO generated drops off markedly as the core depth increases, though the actual spatial kinetics of generation do not bear the linear relationship that superoxide appears to (see inset graph, figure 8.12).

These data illustrate a number of points, firstly, impact stressed potato tuber tissue generates NO, secondly this impact stress induced NO generation is highly localised to the outer 3 mm of the tuber, at the impact site, and finally, as the experimental core is excised from deeper into the tuber the production of NO rapidly tails off indicating a very tight spatial regulation of NO generation. In order to maximise and standardise all subsequent nitric oxide measurements only the outer 3mm of tuber tissue was utilised.

Repeated nitric oxide determination with supermarket tuber material proved just as variable as described in the superoxide quantification experiments (section 8.2.2). It was assumed that the inconsistency stemmed from the same problems – variation in tuber age and potential pre-laboratory impact damage.

# 8.2.4 $O_2^{-1}$ and NO generation in different potato varieties

As stated in sections 8.2.3 and 8.2.4, repeated time course experiments produced variable and less consistent superoxide anion and nitric oxide generation profiles. Likely probable causes of this inconsistency included, tuber age and impact damage prior to arriving at the laboratory. Consequently, it was deemed necessary to acquire potatoes from a more consistent source. The potato research council out-station at Sutton Bridge, Lincolnshire were able to supply a variety of hand collected potatoes that had a known history. Each of these potato varieties were known to have differing degrees of susceptibility to impact damage and consequently provided an excellent opportunity to investigate superoxide and nitric oxide production from impact and control sites within

tubers with varying susceptibility to impact damage. S. Johnson had tested a number of potato varieties for their susceptibility to impact damage, the data are presented below. The arrow diagram below graphically depicts a linear scale of impact susceptibility calculated by the propensity of a tuber to form a bruise (depth of colour and size) in response to impact stress. This index of impact susceptibility was carried out by S. Johnson at the University of Durham.

1------ 2------ 3------ 4------ 5------ 6------ 7------ 8------- 9 \_\_\_\_\_\_

Tuber varietySusceptibility IndexMaris Piper2King Edward4Saturna8Russet Burbank9

Increasing susceptibility to impact damage

Each of these varieties was tested independently for the generation of the superoxide anion (see figures 8.13 graphs A-D) and the nitric oxide molecule (see figure 8.14 A and B) over the six hours after an impact stress had been imposed (see section 8.1.1 for experimental design).

Comparisons of the four superoxide time courses reveal that the overall duplicate measurements of superoxide generation were more stable and less variable using tubers with a known history (compare error bars between figures 8.12 and 8.13 A-D). These data are presumably reflect the greater consistency in growth conditions, storage and impact status of the potatoes from Sutton Bridge. Note also the consistency of superoxide generation at the control sites during all four time courses, generation remained relatively stable over the whole assay period, regardless of which potato variety is being tested (<1.5 nMoles  $O_2$ /g/minute). It was concluded from these observations that the control site is in fact generating a 'background' level of the superoxide anion, that is assumed to be produced as a response to the stress of excising the core for analysis.

Figure 8.13 (A and B) graphically presents the time course experiments for the impact resistant varieties (King Edward and Maris Piper). They both exhibited very little difference between the control and impact sites; and in fact, if we discount the King Edward four hour time point, the impact sites of both varieties stay below the control site superoxide level found in all varieties of <1.5 nMoles  $O_2^{-7}$ g/minute. Overall, no statistically



# Figure 8.13

# Time course of superoxide anion production from impact stressed tuber tissue of different varieties.

Experimental design is described in section 8.2.1.1 and the final assay is presented in section 4.3.5. The experiments were carried out with Maris Piper (graph A), King Edwards (graph B), Russet Burbank (graph C) and Saturna (graph D) potato varieties. The data from the control site and the impact site were statistically assessed (using GraphPad Prism software) to evaluate if there was any significant difference between the two data sets; this is shown below each graph. P-value summary key; ns=no significant difference; \*=significant difference;

significant difference was found between the impact and control sites in either King Edward or Maris Piper impact resistant potato varieties.

The impact susceptible varieties exhibited a very different superoxide generation response than the impact resistant varieties. Figure 8.13 (C) graphically depicts the data for superoxide anion generation over a 6 hour period, post impact damage at the impact and control sites for the impact susceptible potato variety, Russet Burbank. It is immediately obvious that superoxide generation levels at the control site are generally low and relatively stable over the six-hour period, they could be possibly described as, 'basal' and probably arise because of tissue damage caused during core excision (see above and section 8.2.2). However, the impact site exhibits a massive and concerted increase in the production and liberation of the superoxide anion into the extracellular medium, peaking at 5 times the quantity produced at the control site after approximately 4.5 hours. Statistically the production of the superoxide anion at the impact site is significantly different from production of the superoxide anion at the control site, thus reflecting the spatial regulation of superoxide production. Similarly, the other impact susceptible variety, Saturna (figure 8.13, graph D) exhibited a statistically significant difference in superoxide anion generation between the control and impact sites. Most interestingly, the superoxide generation response in both impact susceptible varieties exhibited a distinct biphasic nature, with a peak in generation at approximately 2 hours followed later by a greater generation of the superoxide anion at approximately 4-5 hours, this phenomenon can be seen very clearly in figure 8.13, graph D.

Nitric oxide measurements were generally more variable, even using tubers with a known history. Figure 8.14, graph A, shows the data for a NO time course of the impact resistant Maris Piper potato variety. The control site showed a relatively stable 'background' level of nitric oxide generation that was attributed to the stress of core excision (<0.6 nMoles NO/g/minute). The NO generated at the impact site was significantly greater than the control site, thereby re-confirming that in different areas of the tuber nitric oxide is generated at different rates (i.e.: spatial regulation). Furthermore the production of NO by the impact site appears broadly biphasic peaking initially at approximately 1.5-hours and again at 4-hours. No significant difference was found between the control and impact sites of the impact susceptible variety, Russet Burbank, in fact inspection of the data (figure 8.14, B) reveals that the control site appears to match the NO generation profile observed at the impact site.





Figure 8.14

# Time course of nitric oxide production from impact stressed tuber tissue of different varieties.

### 8.3 Conclusions

The detection and quantification of the superoxide anion and nitric oxide molecule in the extracellular environment and generated by impact stressed tuber tissue using the assay systems and experimental design described here have been successfully confirmed. Although both assay systems were originally designed with the intention of measuring  $O_2^-$  and NO generated from stressed cell suspension cultures they have proved robust and readily adaptable to the direct quantification of these molecules from more physiologically relevant material.

Previous reports have identified mechanical stress as a trigger for the production of AOS, invariably all of these reports have indirectly measured AOS, using the relatively ubiquitous  $H_2O_2$  [Jaffe *et al.* 1985; Legendre *et al.* 1993; Yahraus *et al.* 1995]. Work at the University of Durham using an tuber-impact model had inferred the participation of the hydroxyl radical during the potato response to impact stress [Croy *et al.* 1998]. The study presented here is the first direct confirmation of the production of the  $O_2H/O_2$  radical pair in response to impact stress. Similarly, nitric oxide has very recently been implicated in the biotic stress response [Delledonne *et al.* 1998; Durner *et al.* 1998] this is the first direct confirmation of NO from potato tuber tissue.

It was obvious that the impact site of impact susceptible varieties generated much more superoxide than impact resistant varieties, for instance superoxide production from the Russet Burbank variety peaked at >4 nMoles  $O_2$ /g/minute, whereas the impact resistant varieties did not rise above 1.75 nMoles  $O_2$ /g/minute. The greater generation of superoxide by impact susceptible potato varieties was in agreement with data generated from oxyblot analysis carried out by Croy *et al.* [1998]. Impact resistant varieties showed a distinctly different response with little or no discernible generation of superoxide. It seems highly probable that the impact susceptibility (i.e.: the propensity to bruise) is directly linked to the production of AOS.

Another distinctive feature of the superoxide generation from impact susceptible varieties was a distinct biphasic generation starting with an early burst (1-2 hours) followed by a later and greater production (4-5 hours). Interestingly this biphasic generation of the superoxide anion has essentially the same kinetics of AOS generated by plant cells in response to a bacterial pathogen [Wojtaszek 1997]. The initial production is thought to be a non-specific and general stress response, the latter production is considered specific to incompatible interactions [Levine *et al.* 1994]. This opens up the intriguing possibility that the impact susceptible tuber is responding to impact stress in essentially the same manner as it would a pathogen and asks the question; do the two disparate stresses

share the same recognition pathway? Similar theories of shared signal transduction pathways have been aired previously [Jongsma *et al.* 1994; Baron & Zambryski 1995].

Unfortunately measurements of nitric oxide generation were not as consistent as would have been hoped and compared to the superoxide assay much more variability was evident. However, significantly elevated levels of NO were seen at the impact site of an impact resistant variety. Many reports concerning the actions of NO, detail the synergistic interplay of the superoxide anion and the nitric oxide molecule to form the peroxynitrite radical (OONO<sup>-</sup>) [Delledonne *et al.* 1998; Durner *et al.* 1998]. However, it was not possible to directly measure 'OONO<sup>-</sup> generation in this research, but its formation is presumed to occur because both reactants are present in the extracellular environment. The exact roles that NO and 'ONOO<sup>-</sup> actual play is not yet clear though this series of experiments will provide a good starting point for further investigation.

### **SECTION 9**

### **DISCUSSION & FURTHER WORK**

Vacuum infiltration was successfully utilised to selectively extract apoplastic polypeptides without contamination from the intracellular environment. Many previous workers [e.g.: llett 1998] have utilised VI to investigate individual apoplastic components, but to the authors' knowledge, none have utilised the technique in such a wide-ranging and diverse study of the apoplast.

Direct N-terminal sequencing of SDS-PAGE separated apoplastic polypeptides did provide some sequence data but overall the vast majority of apoplastic polypeptides that underwent sequencing appeared to be N-terminally blocked. Interestingly, in animal systems proteins that are exposed to AOS can undergo modification to their N-termini that ultimately results in blockage [Burcham & Kuham 1997] and in plants the apoplast is a proposed site of AOS generation and release during stress and growth [Lane *et al.* 1992; Wojtaszek 1997; Fry 1998]. The constant exposure of apoplastic proteins to AOS may explain the unusually high level of N-terminal blockage experienced and if correct would also explain why all of the attempted sequencing of apoplastic polypeptides from abiotically stressed plants failed. Several methods do exist for the deblocking of N-terminally blocked polypeptides and may be of use for the sequencing of apoplastic polypeptides.

However, some polypeptides did generate sequence data, which could be utilised for homology searching. Two apoplastic polypeptides (SICF2 and SICF3) were shown to exhibit a degree of homology with cell wall/carbohydrate related enzymes already known to be present in the apoplast [see Fry 1991] and a 27 kDa thaumatin-like protein from Diospyros texana [Vu & Huynh 1994] had some homology to an apoplastic polypeptide of ~26 kDa. Most strikingly however were the number of potential ABP's that were successfully sequenced, high homologies were found to germin-like proteins and ABP's (ITL5, SICF8 and 9) from peach [Ohimya *et al.* 1998].

The putative site of auxin action is thought to be the apoplast [Brown & Jones 1994] although the vast majority of the most well characterised ABP, ABP1 from maize, is resident in the endoplasmic recticulm [Campos *et al.* 1995]. However, recently, circumstantial evidence has accumulated that ABP's are indeed apoplastic, for instance, Tsurusaki *et al.* [1997] have shown that the concentration of endogenous auxin is higher in the apoplast than the symplast and Jones & Herman [1993] and Deikmann *et al.* [1995] have both provided microscopy evidence. Consequently, ICF was subjected to

auxin affinity chromatography. Three potential auxin binding polypeptides were identified by SDS-PAGE analysis and one of these polypeptides may have been sequenced earlier and proved to be a putative ABP (SICF9 ~20.3 kDa). Recently, Shimomura *et al.* [1999] have identified a 19.6 kDa homologue of maize ABP1 that is secreted from tobacco leaf discs into the surrounding medium, this polypeptide and SICF9 may be the same. With two independent pieces of evidence localising a tobacco leaf ABP of ~20 ±1 kDa, to the apoplast a strong suggestion that the active site of ABP's is indeed the apoplast can be proffered.

ITL5 was a 16.5 kDa apoplastic polypeptide that shared 100% homologies with an ABP and a germin-like protein. Some germin-like proteins have been identified as containing auxin responsive elements [Lane *et al.* 1992] and have been tightly linked with cell wall elongation [Lane *et al.* 1992; Ohimya *et al.* 1993]. ITL5 was present only in the immature tobacco leaf tissue and may therefore play roles in cell wall elongation during leaf growth.

The approach of utilising VI to recover ABP's present in the apoplast circumvents a major problem associated with the more commonly employed whole tissue studies, that invariably suffer from the problem that >99% of ABP's are typically present in the endoplasmic reticulum [Campos *et al.* 1995]. The use of VI in the study of ABP's is therefore very important precisely because such low quantities of ABP present in the apoplast are difficult to distinguish from ABP resident in the intracellular environment. Thus, this approach of sampling only the compartment in which ABP's are proposed to be active is extremely novel and worthy of further work.

Early in the project it was identified that 2D electrophoresis would be a useful technique to improve the separation of apoplastic polypeptides extracted using VI. The hope was that smaller changes to the apoplastic complement of polypeptides could be visualised. All the methods detailed within this thesis failed primarily because of the nature of the material. The ionic environment apoplastic proteins and polypeptides inhabit is ionically rich and unfortunately the first dimension of 2D electrophoresis requires that proteins and polypeptides were present in a buffer and gel that has an extremely low ionic strength. This change in the ionic environment coupled with the exposure of the polypeptides to a strong electrical field is assumed to have contributed greatly to the observed and consistent electro-precipitation. A further potential complication associated with ICF samples is varied and heterogeneous nature of other components that are unwittingly extracted alongside apoplastic proteins and polypeptides during VI. Molecules and compounds with the potential to be problematic include highly heterogeneous carbohydrates, phenolic and polyphenolic compounds [Fry 1991; Dawes *et al.* 1994; Dietz *et al.* 1997]. Additionally, the glycoprotein modifications present on all apoplastic

proteins and polypeptides also show evidence of charge heterogeneity [Dawes *et al.* 1994; Amoresano *et al.* 1999] which may explain the smearing of 2D separated polypeptides observed in the ESA Investigator<sup>™</sup> gels. Due to these problems 2D electrophoresis of ICF that attained a suitable standard could not be achieved during this project.

The apoplast has been determined to be a site of production and release of AOS in response to stress and growth [Lane *et al.* 1992; Wojtaszek 1997; Fry 1998]. A range of anti-AOS defences that include the enzymes peroxidase and superoxide dismutase, are known to be present in the apoplast [Fry 1991; Schinkel *et al.* 1998] and are presumed to control the formation of apoplastic AOS.

Native-PAGE in gel enzyme activity staining was successfully utilised to analyse ICF extracted using VI. These techniques revealed 4 ICF-POD's and 5 ICF-SOD's. All of the ICF-POD's separated using native-PAGE contained the metal ion, iron; this was distinct from their intracellular counterparts, all of which appeared to contain the selenium ion. ICF-POD 1 was potentially identified as a dimer with a sub-unit mass of ~29 kDa, a similar sized basic polypeptide with POD enzyme activity has been isolated from the apoplast of pine needles by Slazer & Hager [1993] and Otter & Polle [1997].

ICF-SOD 1 was 20.2 kDa CuZn-SOD with a high isoelectric point, an apoplastic SOD enzyme with a similar mass and isoelectric point has been characterised recently from pine needles [Schinkel *et al.* 1998]. A putative Mn-SOD (ICF-SOD 4) was identified in the tobacco leaf apoplast and potentially is only the second apoplastic plant Mn-SOD ever identified [Yamahara *et al.* 1999]. The remaining ICF-SOD's were Fe-containing SOD's (ICF-SOD's 2 and 3) and to the authors knowledge they are the first apoplastic iron containing SOD's described in plants. An extracellular Fe-SOD has very recently been discovered in bacteria [Shirkey *et al.* 2000] but none have been currently located to the apoplast/extracellular matrix in higher organisms. Coincidentally, the extracellular Fe-SOD from bacteria has almost the same molecular mass as the apoplastic Fe-SOD, ICF-SOD 3 identified within this study.

Very interestingly native-PAGE revealed that ICF-POD's 1 and 2 co-migrated with ICF-SOD 4 which indicated two possibilities, either, these proteins have both SOD and POD enzyme activities, or alternatively they may be independent enzymes with similar separation characteristics in this native-PAGE system. The final explanation proffered for the co-migration in the native-PAGE system is that they were in fact linked in some manner. Johansson *et al.* [1999] has described such an interaction of POD and SOD enzymes via carbohydrate moieties in the extracellular matrix of animals. An interaction between these two enzymes may potentially streamline the process of AOS detoxification in the apoplast.

In the study of the complement of apoplastic, POD's and SOD's VI appears to have yielded unique results. It is possible that the novelty of application of VI coupled with native-PAGE separation and in gel assays to identify individual proteins and polypeptides that are SOD or POD active has yielded such new observations. A similar approach by Schinkel *et al.* [1998] using pine needles has also generated unique insights into apoplastic anti-AOS enzymes. The more commonly employed experimental procedure of extraction of the particular cellular compartment of interest followed by the use of a spectrophotometric assay of all the active components within that compartment, does not yield information on the activities of individual components. Obviously, this approach is worthy of greater investigation, perhaps to the study of other stress related apoplastic enzymes.

Investigation of the generation and release of apoplastic AOS was technically difficult with the leaf-VI system used throughout this research. Consequently, a tuber-impact stress system already in use in Durham was adopted. The system had the distinct advantage that direct quantification of the superoxide anion and nitric oxide molecule in response to a defined stress was possible.

Preliminary studies revealed that superoxide anion production in response to impact stress was spatially regulated. This result concurs with oxyblot data generated by Croy *et al.* [1998]. More detailed impact studies using potatoes with a well known history revealed a distinct difference between impact susceptible and resistant varieties, based upon their propensity to form a bruise in response to the impact stress. Impact resistant varieties did not exhibit superoxide anion production greater than that observed at control sites within the same tuber. Impact susceptible varieties exhibited a distinct biphasic production of the superoxide anion that was extremely reminiscent of the well documented pathogen induced oxidative burst [Wojtaszek 1997; Levine *et al.* 1994]. There is the possibility that both 'stresses' induce a similar oxidative burst in the apoplast [Jongsma *et al.* 1994; Baron & Zambryski 1995].

The presence of apoplastic NO was shown, however, 'bursts' and 'phases' of production in response to impact stress were not as distinct as the measurements of the superoxide anion. Commonly nitric oxide and AOS act synergistically to produce a biological effect [Delledonne *et al.* 1998; Durner *et al.* 1998], via the formation of peroxynitrite from the superoxide anion and nitric oxide. The combination of superoxide and nitric oxide at the site of impact stress would generate the peroxynitrite radical; this radical is an effective anti-microbial agent because of its strong oxidising properties [Mayer & Hemmens 1997].

Thus impact stress and pathogen attack may in fact share the same signalling pathway highlighted by the presence of anti-microbial peroxynitrite [Mayer & Hemmens 1997; Durner *et al.* 1998].

The impact site of impact susceptible potato tubers can be typified by the formation of a bruise and localised cell death [Peters 1996]. Peroxynitrite is involved in the process of apoptosis in animal systems via protein modification to kinases and caspases [So *et al.* 1998; Jianrong *et al.* 1997] and may be playing a similar fundamental biological role in plants. Much more study is required before firm roles can be attributed to peroxynitrite in plants, although both Delledonne *et al.* [1998] and Durner *et al.* [1998] have implicated peroxynitrite in the plant hypersensitive response to pathogen attack.

A potential avenue for further research, which may yield very rapid results, is the investigation of the protein modifications induced by peroxynitrite. The adoption of a 'nitroblot', based on exactly the same principle as the 'oxyblot' [Gow *et al.* 1996] would be a suitable approach for this work. Such an approach would make it possible to investigate which proteins are being modified at the time stress, and ultimately to determine the consequences of modification.

As an adjunct; it is noteworthy that phenolic compounds are intimately involved in the response of potato tubers to impact stress at the impact site, and in particular tyrosine and melanins [Croy *et al.* 1998] essentially form the colouration distinctive of a bruise. These compounds are known antioxidants that are effective against peroxynitrite and AOS [Sandova *et al.* 1997] and furthermore they are extremely indigestible to enzymatic action [Matern *et al.* 1995]. The exact role of phenolic compounds at the impact site is not known, though they would presumably limit pathogen ingress.

## **SECTION 10**

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# **APPENDIX 1**

#### Submitted papers and papers in preparation

#### Doherty, S. & Croy, R.R.D.

Tobacco leaf apoplastic superoxide dismutases and peroxidases. Plant, Cell and Environment Submitted September 2000.

#### Johnson, S.M., Doherty, S. & Croy, R.R.D.

Superoxide radical production in potato tubers: a novel biphasic response to mechanical stress. Planta

Submitted August 2000.

#### Doherty, S. & Croy, R.R.D.

A simple and rapid microtitre plate spectrophotometric assay for superoxide dismutase.

Pre-submission manuscript.

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# Superoxide radical production in potato tubers: a novel biphasic response to mechanical stress

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Abstract. Potato (Solanum tuberosum L.) cultivars differ quantitatively in their ability to produce blue-black melanin pigments in tubers following mechanical Investigations into the events stress. following mechanical stress exerted by direct impact on tuber tissues have shown that an early cellular response is a synthesis of and rapid significant Distinctively this superoxide radicals. burst of radical production displays a reproducible biphasic pattern over time with peaks of accumulation at 2 hours and 5 hours. A concomitant consequence of the generation of these free radicals is elevated levels of oxidatively modified Both radical generation tuber proteins. and protein modification vary between cultivars but both are directly proportional to the amount of melanin pigments produced.

Key words: blackspot bruising mechanical stress - oxidative burst - potato tubers - superoxide radicals

#### Introduction

Mechanical stress is imposed on plant cells by a variety of stimuli and lead to a Early studies on range of responses. mechanical stress used simple physical contact with bean stem tissues to examine growth in cell responses callose (thigmomorphogenesis), deposition and ethylene sythesis (Jaffe 1973, Jaffe et al. 1985). More recently Legendre et al. (1993), Yahraus et al. (1995) and Cazale et al. (1998) exposed sovbean and tobacco cells to mechanical stress by agitation of suspension cultures or by exposing the cells to hypo-osmotic medium to increase cell turgor. Using . able thev were to systems these demonstrate short term generation of hydrogen peroxide. Yahraus et al. (1995) exerted direct pressure on soybean cells on a microscope slide and were able to show histochemically. generation peroxide Other than these examples few studies have been undertaken to investigate the responses to biochemical primary Potato (Solanum mechanical stress. tuberosum) tubers that undergo a physical impact produce melanin-based pigments, leading to a blue-black discoloration of subdermal tissues known as blackspot

Abbreviations: DNPH = 2,4-Dinitrophenylhydrazine; DOPA = 3,4-dihydroxyphenylałanine; PPO = Połyphenol Oxidase; AOS = Active Oxygen Species; XTT = 2,3bis[2-Methoxy-4-nitro-5-sulfophenyl]-2H-tetrazolium-5carboxanilide

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Much of blackspot bruising bruising. occurs during harvesting, transport and storage and is a serious problem for potato growers leading to significant levels of rejection of tuber harvests (Rastovski 1987; Kleinschmidt and Thornton 1991; The melanin 1994 ). Anonymous synthesis pathway in tubers is catalysed in part by the enzyme polyphenol oxidase (PPO) (EC 1.14.18.1). Cellular disruption following mechanical stress leads to decompartmentalisation of the amyloplastlocated PPO which then mixes with monophenolic substrates such as tyrosine. A series of oxygen-dependent reactions red-brown DOPA initially produce intermediates and ultimately lead to the formation of blue-black melanin pigments which can polymerise to water-insoluble complexes (Corsini et al. 1992; Stevens and Davelaar 1996: Friedman 1997). Recent evidence points to the involvement of covalently cross-linked protein in these complexes (Stevens and Davelaar 1996. The synthesis of melanin is 1997). thought to be a defence mechanism in polymerised. insoluble which the complexes form a resistant barrier sealing tuber tissues against the entry and spread of pathogens. The predisposition of tubers to melanin synthesis depends on the growing and storage conditions and exhibits a wide range of genetic variation (O'Leary and Iritani 1969; Hudson 1975; Skrobacki et al. 1989; Dixon 1992; British Potato Council 2000). Transgenic plants expressing anti-PPO gene constructs now form the basis of novel potato varieties highly resistant to bruising and confirm a catalytic involvement of PPO enzymes in this process (Bachem et al. 1994; Krohn et al. 1998).

The oxidative burst, a rapid production of highly active oxygen species (AOS) which include various oxygen free radicals and hydrogen peroxide, is a well characterised response of plant cells to (Bolwell and challenge pathogen Wojtaszek 1997; Lamb and Dixon 1997; The generation of Wojtaszek 1997). superoxide radicals is the first step in the production of AOS Addition of a single electron to molecular oxygen generates the superoxide anion:

 $O_2 + e^{-1} > O_2$ 

and is believed to be catalysed by a plasma-membrane enzvme complex. NADPH-dependent oxidase (Doke 1995), although the possible involvement of an alternative pathway involving a pHactivated cell wall peroxidase has also been suggested (Bolwell et al. 1995). cellular conditions normal Under superoxide radicals are rapidly converted to hydrogen peroxide and oxygen by the enzyme superoxide dismutase. However, during an oxidative burst a vast excess of AOS are produced, which, due to their high reactivity with proteins, lipids and nucleic acids have been proposed as potent local inhibitors of pathogen spread (Tenhaken et al. 1995, Mehdy et al. 1996). AOS have also been implicated in signalling roles and in cross-linking of cell wall components (Brisson et al. 1994; Low and Merida, 1996) and it is likely that they also influence a wide range of metabolic events.

We have used an XTT-based assay to measure directly superoxide radical production in tissues from mechanically stressed potato tubers. The results show that levels of superoxide synthesis vary between potato cultivars which also exhibit differing susceptibilities to blackspot bruising. This synthesis is biphasic with two peaks of superoxide accumulation over a 6 hour period. We have also examined changes in the level of protein modification in mechanically stressed tubers and show a similar correlation with susceptibilities to blackspot bruising. We discuss the possible correlations between superoxide production, protein modification and the synthesis of melanin bruise pigments.

#### Materials and methods

Plant materials. Tubers from selected potato cultivars, specifically grown and carefully harvested to avoid any mechanical stress, were kindly supplied by Dr Adrian Briddon (Sutton Bridge Experimental Unit, Sutton Bridge, Lincolnshire, UK). Other tuber materials were purchased locally from commercial outlets. Five potato cultivars were used - Cara, King Edward, Maris Piper, Russet Burbank and Saturna. All tubers were stored in the dark at 10°C to inhibit greening and sprouting.

Tuber mechanical stress. Tubers were incubated for 48h at 4°C in the dark and then impacted at the

stolon end using a standard falling weight of 240g and a 300mm drop height imparting a force of 0.7J. Impacted and control tubers were incubated for 48h at 26°C to allow maximal synthesis of bruise pigments. Tubers were then eut in quarters at the impact site and the volume of atflected tissue measured as well as a visual estimation of the intensity of pigmentation on a scale of 0 (no discoloration) to 3 (deep blue-black coloration). 30 tubers for each of the five cultivars were used and a mean bruise index calculated based on bruise extent and intensity.

Assay of superoxide radical generation in tuber Tubers of each cultivar were exposed to tissues: the standard mechanical stress and incubated in the dark at 26°C for various periods of time prior to assay for superoxide generation. 5mm3 slices of tuber tissue, excised from the centre of impact sites and from distal control sites were washed thoroughly with distilled water, blotted dry and then incubated at 16°C for 20 minutes in 200µ1 0.6mM XTT in 50mM phosphate buffer, pH 8.2 (Able et al. 1998). The tissue pieces were removed and the assay solution centrifuged (13000g) for 5 minutes. The A450 of the supernatant was taken and converted to jumol superoxide generated using the molar extinction coefficient for the XTT formazan of 23,600 M<sup>+</sup> cm<sup>+</sup> (Sutherland and Learmonth, 1997; communication). personal Μ. Sutherland Superoxide estimations were carried out in duplicate and all assays were replicated.

Estimation of protein modification in tuber tissues: The carbonyl content of oxidatively modified tuber proteins was quantified by the spectrophotometric assay method of Levine et al (1994). Tubers were exposed to standard mechanical stress and incubated for 48h. Tuber proteins were extracted from 150mg of control and impacted tubers in 3ml of 50mM phosphate buffer, pH7.4. The carbonyl groups on extracted proteins were reacted with DNPH and the resulting hydrazone derivatives estimated from the peak absorbance at 355-390mn using a molar extinction coefficient of 22,000 M<sup>-1</sup> cm<sup>-1</sup>. Protein contents of duplicate samples were estimated from the A<sub>280</sub>. Values were expressed as nmol carbonyl mg<sup>-1</sup> tuber protein.

# Results

**Table 1.** Bruise indices representing the relative susceptibility of different cultivars to mechanical stress leading to melanin pigment synthesis. The higher the bruise index indicates a higher susceptibility to mechanical stress. The values for the bruise indices are the mean values for 30 independent estimations for each of the five

Bruise Index
9.2
7.2
7.9
うう
5.0
4.0
3.1

potato of Bruise index estimation Table 1 lists the values cultivars: estimated for the bruise susceptibilities for each of the five experimental cultivars. The cultivars used exhibited a wide range of susceptibilities to bruise development highly susceptible (cv Russet from Burbank) to highly resistant (cv Maris Piper). Our estimates of bruise indices for the cultivars used here are largely comparable with published values using similar methods (Dixon, 1992; British Potato Council, 2000).

Figure 1: Time course (hours post-impact) and levels of superoxide (nunol g<sup>-1</sup> min<sup>-1</sup>) generated in tubers exposed to mechanical stress. The live cultivars tested were A) *Russet Burbank*, B) *Saturna*; C) *Cara*; D) *King Edward*; E) *Maris Piper*. ( $\blacksquare - \blacksquare$  mechanically stressed tissue,  $\blacktriangle - \bigstar$ control tissue). The results shown are the mean values for two independent estimates.

Superoxide production in mechanically stressed tuber tissue: Figures 1A-E show the time course of superoxide generation in tuber tissue slices. All cultivars showed some degree of superoxide production in response to impact. Very significant levels were produced by cv. Russet Burbank and cv. Saturna (Fig. 1A-B) while cv. Maris Piper (Fig. IE) displayed only slightly elevated levels compared with the control values. Most striking was the biphasic nature of the superoxide generation pattern over time with an initial peak response at 1-2 hours post-impact followed by a larger response at 4-5 hours. This pattern was reproducible and apparent in all five cultivars although most obvious in cv. Russet Burbank, cv. Saturna, and cv. Cara (Fig. 1A-C) which also showed the highest The peak values for bruise indices. superoxide generation were plotted against bruise index for the five cultivars to between relationship illustrate the oxidative burst and melanin pigment synthesis (Fig. 2B).

**Table 2.** Tuber protein modification levels of the five potato cultivars following mechanical stress. Results shown are protein carbonyl levels expressed in nmol mg<sup>-1</sup> tuber protein and are the mean values for two estimations corrected for background levels.

Cultivar	Carbonyl Content
Russet Burbank	10.1
Saturna	7.4
Cara	3.3
King Edward	2.9
Maris Piper	2.0

**Figure 2**: Histogram comparisons of cultivar bruise indices (Table 1), levels of protein modification (Table 2) and maximum levels of superoxide generated following mechanical stress (Fig. 1). Results shown are the mean values for two independent estimates corrected for control levels and provides a visual representation of the correlations between A) bruise index (solid bars) and protein modification (cross hatched bars) and B) bruise index (solid bars) and superoxide production (cross hatched bars). Cultivar abbreviations: RB=*Russet Burbank*, SA=Saturna, CA=Cara, KE=King Edward, MP=Maris piper.

Protein modification in mechanically stressed tuber tissue: Table 2 shows the results of assays for oxidatively modified as tuber proteins 48h post-impact indicated by the carbonyl content of extracted tuber proteins. It is immediately obvious that these values (Fig. 2A) closely mirror those for the levels of superoxide generation (Fig. 2B) in that cv. Russet Burbank showed the highest level of modified protein while cv. Maris Piper had the lowest. Furthermore comparison of the bruise indices for the cultivars of protein levels 1) with (Table modification (Fig.2A) and with superoxide generation (Fig.2B) show that there is a between these relationship direct superoxide high Thus variables. protein high to leads generation modification and both are linked to high bruise susceptibility.

Figure 3: Correlation plots between bruise index values, protein modification and levels of superoxide generation. Graph shows the near straight line relationships between the levels of melanin pigment synthesis (bruise index) and both superoxide generation and protein modification. ( $\blacksquare$  —  $\blacksquare$  protein modification,  $\blacktriangle$  —  $\bigstar$  superoxide generation).

#### Discussion

Our observations show that cells in tuber tissues exposed to mechanical stress respond with a rapid production of superoxide radical anions. This oxidative burst is genetically highly variable since different potato cultivars show diverse reponses and quantities of superoxide generated (Figs. 1, 2). The use of nitroblue tetrazolium as a histochemical stain for superoxide radicals indicates that the tuber oxidative burst is restricted to the vicinity of the impact zone in a similar region to melanin pigment synthesis (data not presented). Synthesis of AOS such as hydrogen peroxide and superoxide is a primary response of plant cells to a pathogen attack or to treatment with chemical elicitors but few reports have described this response to a mechanical stimulus. Yahraus et al. (1995) simulated sovbean cell in mechanical stress suspension cultures by hypo-osmotic hydrogen and demonstrated media peroxide synthesis within minutes. They also imposed direct mechanical stress by exerting physical pressure on the cells under a microscope slide and were able to production demonstrate peroxide histochemically after 3 minutes. Legendre et al. (1993) and Cazalé et al. (1998, 1999) used mixing of cell suspensions to impose mechanical stress and showed an oxidative burst of peroxide synthesis. In contrast to these studies, we have demonstrated over a much longer time-scale, the direct generation of superoxide radicals in differentiated plant tissues as a response to mechanical stress. This was most evident in Russet Burbank, Saturna and Cara the cultivars which also exhibited the highest susceptibility to mechanical stress ie, the highest bruise indices (Table 1, Fig. 2B). A unique feature of this superoxide synthesis was the biphasic pattern of generation with peak levels detectable at 1-2 hours and 4-5 hours post-impact. This pattern is reminiscent of the biphasic oxidative burst reported by Baker and Orlandi (1995) in which cultured cells treated with elicitor displayed two peaks of hydrogen peroxide generation over a Phase I was period of 6-8 hours. suggested to be a non-specific biological response to stress while phase 2 was determined to be a specific interaction between the pathogen hrp complex and the plant receptors leading to hypersensitive cell death. In the present tuber system we see an initial peak which probably corresponds to the non-specific response. However, our second peak is potentially of much greater interest. The observation that a second burst arises in mechanically stressed tissues in the absence of elicitors or pathogens suggests the possible role of a cell receptor for detecting mechanical stimuli and an activation process. This is the first demonstration of what may be a specific response to mechanical stress and while its nature remains unclear it perhaps activation transcriptional or reflects initiation of signalling pathways by an unidentified mechanoreceptor as proposed by Cazalé et al. (1999). Jaffe et al. (1985). in a study of thigmomorphogenesis in bean stems, reported a minor transient deposition of callose 1.5 hours after mechanical stress in the form of gentle abrasion, followed by a major peak of hours. after 6 callose deposition Interestingly the time course for this response is nearly identical to the superoxide generation response reported here and both studies investigate responses tissues to differentiated plant in mechanical stress and it is tempting to speculate that both phenomena are similar mechanical initiated through receptors and signalling pathways.

The observation that the cultivars generating a large oxidative burst were also those exhibiting high melanin synthesis posed the question as to the relationship between the two processes (Fig. 2B). The correlation plot shown in figure 3 shows that there is a direct quantitative relationship. The near straight superoxide between obtained line generated and bruise indices provides of this proof direct quantitative relationship. Whether this is simply a fortuitous relationship or if there is an interaction between the two processes is Direct unresolved question. an involvement of oxygen free radicals in the enhancement of the melanin synthetic pathway or the PPO reactions has not yet been demonstrated. However melanin and related complexes are known to be free radical scavengers and melanin radicals have been demonstrated in animal pigments under conditions which elevate free radicals (Qu et al. 2000), so it seems plausible that radical production and melanin synthesis may be causally related.

One of the consequences of production is the oxidative radical modification of proteins in which carbonyl introduced are through groups modification of lysine, proline and arginine residues (Stadtman 1993). Such protein modifications are well documented in animal systems but this report is the first demonstration of the effect occurring as an outcome of mechanical stress in Since oxidative modification of plants. tuber proteins arises through interaction with the oxygen free radicals it is not levels of protein unexpected that modification produced a similar straight line relationship with bruise indices (Fig.3). Clearly the level of protein modification is directly proportional to levels of superoxide generated and independent provides therefore corroboration of the levels of radical production.

We present here some novel observations on the responses of plant tissues to mechanical stress based on a convenient experimental system. Further work is needed with this system to confirm and elucidate a causal relationship between radical production and melanin synthesis. Additionally questions remain as to how radicals, which are generated extracellularly, gain access to the largely intracellular tuber proteins in order to The nature and effect modification. function of the detection system for the remains `to be mechanical stress elucidated.

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# A simple and rapid microtitre plate spectrophotometric assay for superoxide dismutase

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Direct assays for superoxide dismutase activity are time consuming, necessitate expensive equipment and high levels of expertise. In contrast, indirect assays are simple, speedy and inexpensive. Most indirect assays are two stage, with an initial generation of the superoxide anion and subsequent reduction of an indicator substance by the anion. Superoxide dismutase in such a system suppresses the reduction of the indicator thereby permitting measurement of activity. Commonly, an enzymatic superoxide generating system is utilised, such as the widely employed xanthine oxidase system. However, it has been noted that this system suffers not only from instability caused by rapid loss of xanthine oxidase activity but also limitations because of enzyme-indicator auto-reduction. Often a convenient indicator is nitroblue tetrazolium, but this too can introduce more acute problems; its reduction is not O2-specific and the resulting mono- and di-formazan dyes are sparingly soluble in aqueous solutions. Consequently, optical density measurements are less than perfect. We have developed an assay utilising a stable non-enzymatic O2-generating system based on the NADHdependent oxidation of PMS with concurrent production of O2 which reduces a new tetrazolium compound, XTT, to form a soluble dye with an absorbance maxima of approximately 450nm.

Superoxide dismutase ( $O_2 : O_2^-$  oxidoreducatse, E.C. 1.15.1.1.) (SOD)<sup>2</sup> catalyses the dismutation of the superoxide anion (SOA) into molecular oxygen and hydrogen peroxide (1). SOA's can originate from a multitude of biological processes, for instance, redox reactions in mitochondria and chloroplast organelles (2, 3), the auto-oxidation of flavins (4), hydroquinones (5) and catecholamines (6), the enzymatic actions of flavin oxidase. xanthine oxidase. aldehvde dehydrogenases (7) or dioxygenases (8). Furthermore, SOA's are produced in a controlled manner by neutrophils and macrophage during phagocytosis (9) and in plant-pathogen interactions which ultimately produce the characteristic hypersensitive response (10). The SOA has a relatively low toxicity primarily due to its very short half-life (2). However, molecular interaction with hydrogen peroxide, especially in the presence of trace metals (eg: Fe2+), can lead to the formation of the relatively long-lived and highly toxic hydroxyl radical (OH). Consequently, SOD has been frequently implicated in playing a primary defensive role against oxidative injury.

Direct assays of SOA and thus consequent measurement of SOD activity typically require high levels of expertise and expensive equipment. They rely on two distinguishing physical characteristics of the SOA, electron paramagnetic resonance (EPR) and UV absorption (11). The latter method necessitates time consuming and extensive sample cleanup prior to because of the inherently high UV analysis, absorbance typical of biological samples. EPR permits direct monitoring of SOA's because the molecule contains one unpaired electron (12). Unfortunately, the resonance frequency can only be distinguished from high background values at extremely low temperatures (4). Consequently, it is unsuitable for routine analysis.

A number of indirect SOD assays have been developed all of which detect reduction of an indicator by artificially generated SOA's. One of the most common ways to artificially produce SOA's utilises the enzyme xanthine oxidase (XO). XO-SOA generation can then be coupled to indicators such as ferricytochrome c or nitroblue tetrazolium (NBT) (13). The XO-SOA generating system suffers from several drawbacks. XO rapidly loses activity after preparation, consequently, later measurements in an assay series have lower SOA formation with a concurrent increase in the level of error. In addition, XO alone can cause indicator reduction when using either cytochrome c (14) or NBT (15) as the indicator, this process does not require the SOA and thus increases false positive error. Futhermore using NBT as an indicator also has inherent problems, the reduced forms of NBT, monoand di-formazan dyes are sparingly soluble in water, an unfavorable property for accurate optical density measurements (15).

A non-enzymatic system of SOA production utilising aerobic mixtures of NADH and phenazine methosulphate (PMS) has been reported (16). This system when tested with NBT removes the inherent problems associated with XO-based SOA generation but still ultimately produces insoluble products. XTT (2,3-bis-[2-Methoxy-4-nitro-5-sulphophenyl]-2H-

tetrazolium-5-carboxanilide) has been employed as a more promising indicator of SOA production and consequently SOD activity (17). We describe here the development of a SOD assay that combines these two very desirable features. The assay has been designed for rapidity of sample throughput utilising a 96 well microtitre plate format. SOA production is achieved using the PMS-NADH system thereby obviating the

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<sup>&</sup>lt;sup>2</sup> Abbreviations used; EPR, Electron paramagnetic resonance; 'OH hydroxyl radical; NBT, nitroblue tetrazolium; PMS, phenazine methosulphate; SOA/O<sub>2</sub>', superoxide anion; SOD, superoxide dismutase; NO, xanthine oxidase; NTT, (2,3-bis-[2-Methoxy-4-nitro-5-sulphophenyl]-2H-tetrazolium-5-carboxanilide) sodium salt.

problems associated with the enzyme XO. The resulting coloured dye has an absorption maxima of 450 nm and is highly stable, soluble and permits measurements to be taken at any point from 2 minutes to 18 hours after addition of sample.

#### MATERIALS AND METHODS Chemicals

SOD (E.C. 1.15.1.1.; 3500 units/mg from bovine erythrocytes),  $\beta$ -NADH, NBT, XTT, EDTA, and PMS were purchased from Sigma Chemical Company (St. Louis, MO, USA) and were of analytical grade. All aqueous solutions were prepared with 18 Mohm/cm ultrapure water from an Elgastat UHQPs system (USF Elga Ltd., High Wycombe, UK).

#### **XTT-SOD** microtitre plate assay

For design and optimisation of the assay, 200µl of 50mM phosphate buffer, pH 7.8 containing, 0.1mM EDTA,  $30\mu$ M XTT and  $98\mu$ M NADH was pipetted into the wells of a microtitre plate.  $25\mu$ l SOD containing a specified number of units in 50mM phosphate buffer, pH 7.8 or 50mM phosphate buffer pH 7.8 (buffer blank) was added. The reaction was initiated by the addition of  $25\mu$ l of  $33\mu$ M PMS. The microtitre plate was agitated for 5 seconds. The optical densities of each well were read at 450nm at 30 second intervals for 10 minutes and at 18 hours after initiation, using a Dynatech MR-5000 plate reader (Dynatech Laboratories, UK).

#### Preparation of blowfly muscle mitochondria

Control flies were grown at a constant 24°C. Flies were heat-shocked by incubating batches at 37°C for 40 minutes. After heat treatment they were returned to a constant 24°C chamber for 4 hours to permit the heat-shock responses to develop. Blowfly muscle mitochondria were isolated as described previously (18). Briefly, thoraces were collected and flight muscles removed and disrupted using a mortar and pestle. Mitochondria were centrifuged at 3,600 rpm (RCF = 1500g) in a Beckman Avanti 30 (F2402 rotor). The mitochondrial pellet was resuspended in 0.15M KCl, 1mM EDTA, 10mM Tris and stored frozen at  $-80^{\circ}$ C.

#### **Protein Assays**

The protein contents of mitochondrial and other samples were determined by the Bradford assay using serial dilutions of standard bovine serum albumin (Bradford, 1976).

#### Data Analysis

GraphPad Prism<sup>®</sup> software version 2.1 (GraphPad Inc.) was used to produce standard curves, perform the regression analysis, quantify SOD levels, and calculate standard deviations (SD).

#### **RESULTS & DISCUSSION**

Principle of the assay



# Optimising O<sub>2</sub> generation & assay duration

The addition of PMS to the reaction mixture initiates the generation of SOA's and consequently XTT reduction and coloured product generation. To establish the efficacy of XTT reduction by O2 generated by the PMS-NADH system and to ascertain suitable assay duration a series of PMS а concentrations were kinetically assayed at 450nm against a buffer blank lacking PMS. Figure 1 illustrates the principle of the assay in which XTT is reduced in a PMS concentration dependent manner. At all PMS concentrations tested absorbance change was linear over a 3 minute assay period (>0.9965 (p<0.0001)). However, continued monitoring beyond 3 minutes for 3.2µM PMS and beyond 4 minutes for 2.8µM PMS, showed that the absorbance change quickly levelled out, indicating that the XTT concentration had been diminished below the level required to react with the constant stream of  $O_2^-$  ions produced from the PMS-NADH generating system. Endpoint measurements beyond 4 minutes and up to 18 hours illustrated that the dye formed during the first 4 minutes remains stable because readings are virtually identical (data not shown).



Figure 1 : The effect of PMS concentration on XTT reduction. Assay conditions were the same as described in Materials & Methods except that the PMS concentration/assay was adjusted. The reaction was initiated with 50 $\mu$ l of varying concentrations of PMS in 50mM phosphate buffer, pH 7.8. OD<sub>450</sub> was measured every 30 seconds over 3 minutes. The final concentration of PMS/assay is displayed in the box. Data are means of  $\pm$  SD of four measurements.

As would be expected maximal absorbance change occurs at the highest PMS concentration,  $3.2\mu$ M. Unfortunately at this concentration the reaction is extremely rapid and it is difficult to attain an accurate initial 'zero time' value (see figure 1). At  $2.8\mu$ M PMS, XTT reduction produced an average absorbance change per minute of approximately 0.1 OD<sub>450</sub> units.

The concentration of PMS was therefore standardised to  $2.8\mu$ M/assay with absorbance readings taken every 60 seconds over a 10 minute period and again at 18 hours. These conditions were used throughout all optimisation experiments. After 5 minutes optical density measurements did not alter over an 18-hour period. Consequently, XTT reduction by O<sub>2</sub> generated by the non-enzymatic PMS-NADH system under these conditions is extremely stable and reproducible.

## **Optimising the XTT concentration**

Reduction of all of the  $O_2$  ions generated by the optimised PMS-NADH system is a necessary prerequisite for accurate measurement of the activity of SOD to dismutate SOA's. XTT concentration is therefore critical to kinetic measurements and requires standardisation to ensure that an XTT-saturation state is achieved during the assay period. This was assessed by performing the assay over a range of XTT concentrations; figure 2 depicts this data.



Figure 2 : Effect of XTT concentration on the reaction rate (OD<sub>450</sub> change/minute). Assay conditions; 50mM phosphate buffer, pH 7.8 containing,  $80\mu$ M EDTA,  $78.4\mu$ M NADH and  $6-150\mu$ M XTT (final assay concentrations). The reaction was initiated with a final PMS concentration of 2.8 $\mu$ M/assay. An endpoint OD<sub>450</sub> was taken at 2 minutes from which OD change/minute was calculated. The final concentration of XTT/assay is displayed on the abscissa. Data are means of  $\pm$  SD of four measurements.

Absorbance change per minute continuously increased with higher concentrations of XTT up to 120 $\mu$ M/assay. At this concentration and above, XTT reduction levelled off. In effect, at this concentration, all of the O<sub>2</sub> ions produced by the PMS-NADH system were instantaneously reducing the XTT producing a maximal absorbance change. When the PMS concentration was set at 2.8 $\mu$ M/assay and the XTT concentration at 120 $\mu$ M/assay, the absorbance change at 450nm over 2 minutes was linear ( $r^2 = 0.9980$ , p<0.0001) and therefore was optimal for accurate kinetic measurements of SOD activity.

#### Defining optimal assay pH

It has been noted that the PMS-NADH system has a tendency to produce hydrogen peroxide at acidic pH and to preferentially generate O<sub>2</sub> ions at more alkaline pH (20). We therefore assessed how pH influenced the PMS-NADH SOA-driven reduction of XTT. Figure 3 illustrates that XTT reduction is in fact pH-dependent, the reaction rate rises sharply from  $<0.09 \text{ OD}_{450}$  change/minute at pH 6.5 to  $>0.12 \text{ OD}_{450}$ change/minute in the pH range 7.5-8.7. The lower optical density readings exhibited below pH 8.0 are probably attributable to the PMS-NADH system producing a variable mixture of hydrogen peroxide and SOA species. Between pH 7.5 and 8.7. SOA production is much more consistent as evidenced by the relatively stable readings depicted in figure 3. At pH >8.7 XTT reduction drops markedly ( $<0.07 \text{ OD}_{450}$ ) change/minute). This may arise because the PMS-NADH system cannot effectively produce O2 ions at pH values >8.7, or because XTT is not effectively reduced at this pH. Therefore, for routine analyses we standardised the assay buffer to pH 8.2, although any point within the pH range 7.5-8.7 is suitable.



Figure 3 : pH-dependence of XTT reduction.

Assay conditions; 50mM phosphate buffer, pH in the range 6.5-8.9, containing,  $80\mu$ M EDTA, 78.4 $\mu$ M NADH and 120 $\mu$ M XTT (final assay concentrations). The reaction was initiated with a final PMS concentration of 2.8 $\mu$ M/assay. An endpoint OD<sub>450</sub> was taken at 2 minutes from which the OD change/minute was calculated. Data are means of  $\pm$  SD of four measurements.

#### **Optimised SOD assay conditons**

The final optimised assay conditions used for both the preparation of the SOD standard curve and for routine sample analysis were: 200µl of 50mM phosphate buffer, pH 8.2, containing 0.1mM EDTA, 98µM NADH, 150µM XTT. 25µl of SOD solution or sample were added and the assay initiated with 25µl of 28µM PMS in 50mM phosphate buffer, pH 8.2. This mixture had final concentrations of 80µM EDTA, 78.4µM NADH, 120µM XTT and 2.8µM PMS. If a microtitre plate reader is not available volumes can simply be scaled up by a factor of four and the measurements made using a spectrophotometer and standard 1-ml plastic cuvettes. Mixing of the reactants can be achieved as with an ordinary cuvette-based assay using parafilm and rapid inversion of the cuvette.

# Production of a SOD standard curve and analysis of unknown SOD samples

A commercial preparation of SOD from bovine erythrocytes, dissolved in 50mM phosphate buffer, pH 8.2 was used to prepare the standard curve. SOD was added to the reaction mixture in a 25 $\mu$ l aliquot in the range 0.01U (2.875 ng) to 100U (28.75  $\mu$ g). Figure 4 illustrates the standard curve. Optical density readings were taken kinetically early after the addition of PMS (<5 minutes), or as endpoint readings (>7 minutes and up to 18 hours). Providing measurements are taken at a standardised time after the addition of PMS any point between 2 minutes and 18 hours provides highly reproducible data and permits inter-assay comparisons to be drawn.



**Figure 4 : SOD inhibition of XTT reduction - standard assay** The figure illustrates the final optimised assay as described in the text.  $OD_{450}$  readings at 7 minutes minus positive control are plotted against the Log. of the units of SOD in the reactions. Data are the means of  $\pm$  SD of three measurements.

We have tested the utility of this assay using control and mitochondria from heat-shocked blow flies. Flight muscle mitochondria were isolated from flies as described previously (20), grown at a control temperature (24°C) and from those undergoing heat shock (36°C) treatment. Heat-shock as with many stresses increases oxidative metabolism, SOD is the only known enzyme that can dismutate O2 radicals before they react with other components to form more reactive and deleterious molecules such as the OH. hydroxyl radical. The activity of mitochondrial SOD is known to be elevated in response to stress (21, 22 & 23). These samples were assayed and the data is presented in Figure 5. As expected the mitochondria isolated from heat-shocked flies showed considerably higher levels of SOD activity confirming the results presented in the literature (21, 22 & 23).



Figure 5 : Assay for SOD in unknown samples

Analysis of unknown mitochondrial samples (see text for details of assay) with endpoint measurements taken at 7 minutes and 18 hours. Data are means of  $\pm$  SD of three measurements.

Optical density readings were taken at 7 minutes and at 18-hours after the addition of PMS to the reaction mixture. There was almost no change in the absorbance readings between the two times, illustrating the stability and longevity of the reduced form of XTT.

The assay described has significant advantages over previously described versions: i) it is large-scale and can accommodate up to 96 samples in a convenient microtitre plate format allowing multiple replicates; ii) it is rapid and simple to setup; iii) it does not suffer from the instability observed in the enzymatic SOA-generating system; iv) it does not suffer from interference from a sparingly soluble dye product; v) it has the added attraction of an extremely stable dye product formation which permits the user to read the assay results anytime between 2 minutes and 18 hours after reaction initiation. This allows rapid reaction rate (kinetic) measurements and reaction endpoint readings to be taken.

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# A simple and quick method to selectively sample large quantities of leaf apoplastic proteins

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#### ABSTRACT

The extracellular compartment of many plant tissues contains pockets of gas confined by the surrounding cells in the tissue and is often referred to as the apoplast. Lining the apoplast is a thin film of moisture that contains many types of proteins including structural- and defence-related proteins and enzymes. The apoplast protein mixture can be isolated largely free from intracellular contamination using vacuum infiltration. This is achieved by submerging the plant tissues in an appropriate buffer, applying a low vacuum which removes gases trapped within the apoplast. A slow release of the vacuum causes the apoplast to flood with buffer. The intercellular fluid (ICF), comprising of the buffer containing dissolved proteins, can be recovered from the apoplast by low speed centrifugation. In order to sample larger quantities of the apoplast proteins for purification and analyses, a large-scale version of this technique has been developed and We have used this technique to optimised. differences in the extracellular investigate complement of proteins in tobacco leaves.

*Keywords*: apoplast; cell wall; intercellular fluid; malate dehydrogenase; SDS-PAGE; tobacco; plant leaves; vacuum infiltration.

Abbreviations: ICF – Intercellular fluid; SDS-PAGE – Sodium dodecyl sulphate polyacrylamide gel electrophoresis; TE – Total extract; VI – Vacuum infiltration; RCF - Relative centrifugal field; RUBISCO – Ribulose bisphosphate carboxylase.

#### INTRODUCTION

The extracellular environment of plant tissues can be viewed as an extension to the cytoplasm but with a unique biochemistry and composition which alters according to changing conditions and stresses. The 'apoplast' is the term used to define the space occupied by pockets of gas transiently entrapped by the aqueous films lining the spaces between cell walls (Canny 1995). Within and exposed to these films of water a variety of proteins are found associated with the cell wall matrix or freely soluble, depending upon the local pH and ionic composition. Examples include peroxidases (Welinder 1991, Mäder 1980, Espelie et al 1986, Fry 1986), carbohydrate- and pectin-related enzymes (see Varner & Taylor 1989 for a review) and a repertoire of hydrolases capable of hydrolysing a wide range of glycosidic linkages present in the cell wall (Hatfield & Nevins 1987) and elsewhere.

Many common abiotic stresses such as, temperature (Marentes et al 1993), salinity (Holland et al 1993). drought (Dhindsa 1991), and wounding (Showalter 1993), alter the content and properties of the cell wall and apoplast, permitting the plant to adapt to the prevailing conditions. Similarly adjustments are evident during biotic stress which establishes initially, a rapid enhancement in the rigidity of the cell wall with a concurrent general reduction in the digestibility of the tissue (Matern et al 1995; Robertson et al 1995; Showalter 1993). This is rapidly followed by secretion of a range of pathogenesis-related proteins (Apel et al 1990; Konbrink et al 1988) and antimicrobial compounds (Dixon 1986) into the apoplast.

Vacuum infiltration (VI) is a relatively simple, vet highly effective strategy, for selectively extracting soluble and ionically-bound proteins from the extracellular compartments. It is the only method that can be used to selectively isolate extracellular components from differentiated plant tissues. Once optimised VI generally causes very little cellular damage and therefore intracellular contamination of such isolates is minimised. While there is a wide range of potentially valuable and interesting proteins and enzymes present in the apoplast the small amounts of material recovered by this technique prevent its wide application for the study of these components. We describe here the development and optimisation of a large-scale version of the Hammond-Kosack (1992) VI technique that typically can yield 30-35 ml of approximately 100µg protein/ml intercellular fluid (ICF) in an 8-hour working day. Using this modified technique we have investigated development-specific changes in the enzyme and protein complement in the apoplast of tobacco leaves.

#### MATERIALS AND METHODS Plant Material

Seeds of tobacco (*Nicotiana tabacum*) cv *SR1* were sown in Levington M3 compost (Levington, UK). Seedlings that had emerged after 14 days were transferred to 2 litre pots and grown in a controlled environment room at  $23/20^{\circ}$ C ( $\pm 2^{\circ}$ C) day/night temperatures. 16h photoperiod of  $100\mu$ E s<sup>-1</sup> m<sup>2</sup> (50W.m<sup>2</sup>) light intensity (cool-white fluorescent tubes and  $70\% \pm 5\%$  relative humidity. Plants were fertilised every two weeks with 1% (w/v) Phostrogen (Phostrogen, UK) and watered once every three days. Unless stated otherwise all plant material used was 8 weeks old.

#### Total soluble protein extraction (TE)

Tobacco leaves (<5 cm and >10 cm length) were excised from plants at 8 weeks old and strips  $(\sim 8-10 \text{ cm}^2)$  were cut from between the major leaf veins using a sharp blade. The total tissue fresh weight was recorded. The tissue strips were placed in an icechilled mortar, with sand and liquid N2 and ground to a fine powder. Extraction buffer (1mM DTT or 1mM DTT, 200 mM NaCl, pH 7.5) was added at a ratio of 1.5ml per gram of tissue. The tissue was further ground until a thick slurry was produced. Clarification of the TE was achieved by an initial 5 minute spin in a bench-top micro centrifuge (RCF=10,000g), followed by ultracentrifugation of the supernatant at 200,000g, for 60 minutes at 4°C. The supernatant was decanted and dialysed for 24 hours at 4°C against dH<sub>2</sub>0 and stored at -20°C or lyophilised.

#### Vacuum infiltration (VI)

VI was carried out in a similar manner to that of Hammond - Kosack (1992). Tobacco leaves (<5 cm and >10 cm length) were removed from 8 week old plants. Strips of leaf tissue (~8-10 cm<sup>2</sup>) were cut from between the major leaf veins using a sharp blade and total tissue weight was recorded. The strips were washed for >20 minutes in two changes of distilled water at 4°C, with occasional mixing to remove soluble proteins from the cut cells. The tissue strips were thoroughly dried using paper towels and arranged on a nylon mesh (1m x 7 cm; mesh size ~1mm<sup>2</sup>) which was then gently rolled to form a 'Swiss-roll' type tube as shown in Figure 1.



Figure 1: Arrangement of leaf strips on the nylon mesh prior to rolling into a 'Swiss-roll' as shown on the right, and secured with an elastic band.

To maintain the roll integrity during manipulations, an elastic band was loosely fixed around the nylon mesh. Several assembled rolls were then placed in chilled infiltration buffer (1 mM DTT or 1mM DTT, 200 mM NaCl, pH 7.5) in a vacuum vessel equipped with a valve suitable for the slow release of the vacuum. The rolls were kept submerged in the infiltration buffer throughout the procedure using a metal grid and a 1kg weight. Evacuation of gases trapped in the apoplast was achieved by applying a standard vacuum (8.5 x 10<sup>4</sup> Pa (Nm<sup>-2</sup>) to the surface of the liquid and gently agitating for 30 seconds. Infiltration of the apoplast with the buffer was then achieved by slowly releasing the vacuum over a 30-second period. After infiltration the tissue rolls were removed from the buffer, wrapped in paper towels to drain off, excess liquid trapped in the nylon mesh. The rolls were then inserted into 25 ml medical sample tubes (25mm diam x 90mm long) with a 3-4 mm hole cut in the bottom. Individual sample tubes were then placed into 50 ml

conical centrifuge tubes (Falcon). The tube assemblies were then centrifuged in a Sigma 204 centrifuge at up to 3000 rpm (RCF=~1500g at R<sub>max=150mm</sub>) for 10 minutes in a swing-out rotor at 4°C. The ICF's collected in the centrifuge tube were usually russetcoloured or almost colourless. ICF's coloured green due to leaked chloroplast material indicated gross contamination with intracellular contents and were immediately discarded. ICF's were further assessed for intracellular contamination by assaying for malate dehydrogenase (MDH). ICF's with an MDH-activity >1% of the level in the leaf total extract were discarded while those with activities <1% were pooled, dialysed against distilled water, and lyophilised.

#### Malate Dehydrogenase and Protein Assays

Malate dehydrogenase (MDH, EC 1.1.1.37) activity was assayed in both total soluble leaf protein and ICF's according to the Biochemica Enzymes Catalogue (Boehringer Mannheim, 1<sup>st</sup> edition, 1968). The reaction mix comprised of 943 $\mu$ l of 0.1M phosphate buffer, pH7.5, 33 $\mu$ l of 2mg/ml sodium oxaloactetate in phosphate buffer, 17 $\mu$ l of 10mg/ml NADH (Na<sup>+</sup>). The reaction was started by adding up to 7 $\mu$ l of either TE or ICF to the reaction cuvette and the decrease in absorbance at 340 nm was measured for 60 seconds. In order to 'quantify' the degree of intracellular contamination of the ICF, TE readings were standarised to 100% of the MDH-activity present in the tissue according to the formula (Hammond-Kosack, 1992): -

 $\frac{\Delta A b^{340} \text{ of ICF } x100}{\Delta A b^{340} \text{ of TE}} = \text{Relative MDH activity of ICF}$ 

The protein contents of samples were assayed using Bradford microtitre plate assays using serial dilutions of a 1mg/ml BSA for calibration (Bradford 1976).

#### **SDS-PAGE & Gel Staining**

The polypeptide composition of ICF and TE samples were analysed by SDS-PAGE performed essentially as described by Laemmli (1970) using 0.75 mm-thick. 12%(w/v) acrylamide gels (acrylamide/bisacrylamide (37.5:1) separating gels). acrylamide vertical slab gels (Bio-Rad Mini-Protean II system). Protein bands were visualised by staining with a silver staining kit (Bio-Rad), following the manufactures instructions. Native-PAGE analysis was performed as described above except samples were not denatured to ensure that the enzymes remained active. Gels were stained for superoxide dismutase (SOD) activity by incubating the gel in a solution of 50 mM Tris-HCl, pH 8.5 containing, 0.98 µM nitroblue tetrazolium, 1.57  $\mu M$  phenazine methosulphate and 5.9  $\mu M$ magnesium chloride with agitation under a strong light. SOD activity is distinguished as clear bands on a dark blue background (Manchenko 1994).

#### RESULTS

# **Optimisation of Vacuum Infiltration**

It was clear from initial experiments that the length of time tissues are exposed to the vacuum and the level of vacuum applied both affect the efficiency of VI as well as the integrity of the cells. This was judged on basis of the observed increase in MDH activity in ICF's from tissues on prolonged exposure to the vacuum or to vacuums higher than  $8.5 \times 10^4$  Pa Consequently. presented).  $(Nm^{-2})$ (data not optimisation of the length of time that tissues can be exposed to this vacuum with minimal intracellular leakage was determined. A typical set of results is shown in figure 2. Exposing tobacco leaf tissues to the vacuum for 60-90 seconds resulted in a stable (background) level of relative MDH-activity in ICF's of between 0.2-0.4% of the MDH-activity present in the TE. Extending the vacuum exposure time to >90 seconds for this tissue, resulted in increasing ICF MDH-activity indicating greater cell disruption and leakage of intracellular components.



**Figure 2**: Change in MDH activity in ICF's with prolonged exposure to the vacuum  $(8.5 \times 10^4 \text{ Pa})$ . Four sets of four tissue rolls were prepared as described in the text. Each set of rolls was subjected to 30, 60, 90 or 120 seconds under vacuum. ICF's were then recovered and the MDH activities assayed. Points plotted are the means of the four results for each time point.

The Swiss-roll arrangement of tissue and nylon mesh affords some protection to the plant material and provides mechanical support during infiltration and centrifugation. However, excessive or prolonged centrifugation of the tissue rolls can cause them to be crushed at the bottom of the sample tube. Deformation of the tissue material inevitably causes cellular damage and gross leakage of intracellular components into the ICF. This was borne out experimentally as



**Figure 3**: Increasing MDH activity in ICF's isolated at increasing centrifugation speeds (RCF). Six sets of four rolls were prepared by infiltration for 30 sec under vacuum as described in the text. ICF's were recovered by centrifuging sets of rolls at one of the following speeds (RCF): 500rpm (42g), 1000rpm (168g), 1500rpm (377g) 2000rpm (671g), 2500rpm (1048g) or 3000rpm (1509g). The points plotted are the means for the four results for each centrifuge speed.

shown in Figure 3. Relative MDH-activity in the ICF steadily increased as the RCF was increased from <50g up to >1500g; at the higher speeds, the supporting mesh tended to distort at the bottom of the sample tube. Consequently, the tissue within the mesh was crushed and the relative MDH-activity was therefore higher in these samples. At higher centrifugation speeds both the volume of ICF recovered and it's protein content were much higher than the final optimised extraction procedure (*data not shown*), however the ICF's were invariably discarded because of obvious intracellular contamination.

Both time of exposure to vacuum and centrifugation speed will influence the quality and quantity of ICF recovered from vacuum infiltrated tissues. For tobacco leaf tissue this large scale VI procedure has now been standarised to a 60-second exposure at a vacuum of  $8.5 \times 10^4$  Pa (Nm<sup>-2</sup>) with recovery of ICF's at an RCF of <200g.

#### **Gel Analysis**

As an example of the application of this modified technique, ICF and TE samples were both prepared from immature (leaves <5 cm in length, from the upper third of the plant) and mature (leaves >10 cm in length, from the bottom third of the plant) leaf tissues. The samples were then subjected to SDS-PAGE gel analysis an example of which is shown in Figure 4.



**Figure 4**: SDS-PAGE analyses of ICF ( $38\mu$ g total protein) and TE ( $60\mu$ g total protein) samples isolated from mature and immature tobacco leaves. (M = size marker proteins).

Examination of the silver-stained gels showed firstly that contamination of ICF samples with intracellular protein was minimal as evidenced by the conspicuous near absence of major intracellular leaf proteins such as RUBISCO, in either ICF sample. Secondly, it was obvious that there were few if any common polypeptides between the respective TE's and their ICF's again indicating the selective enrichment of components from the extracellular compartment. Thirdly, comparison of the analyses of immature and mature ICF's showed several common protein components presumably representing those constantly development. However, required throughout differences were clearly evident - bands specific to either immature or mature tissue were clearly discernable presumably reflecting protein components functioning at specific development stages in that tissue.



**Figure 5**: Native-PAGE analyses of ICF (15, 37 and 75 $\mu$ g total protein) and TE (16 $\mu$ g total protein) samples stained for the presence of SOD.

We have also investigated the distribution of the enzyme superoxide dismutase (SOD) between extracellular and intracellular compartments. Figure 5 shows *in situ* staining of a native gel containing a total protein extract and ICF from tobacco leaves. Interestingly the apoplast contains a unique complement of basic SOD isoforms which are not detectable the total extract.

#### **DISCUSSION**

It is well documented that small amounts of soluble or weakly bound proteins can be selectively sampled from the apoplast using VI followed by gentle centrifugation (Hammond-Kosack 1992; Rohringer et al 1983: Söding 1939; Terry & Bonner 1980). The VI procedure described here is as effective as these previous techniques at selectively enriching apoplastic material, according to contaminating enzyme assays (MDH) and gel analyses. However the scale of isolation has been increased by at least an order of magnitude such that enough ICF can now be conveniently isolated to make protein purification and proteomic analyses realistic possibilities. The amounts of protein isolated by this modified technique are in the milligram quantities, easily enough to conduct one and two dimensional PAGE analyses combined with N-terminal amino acid sequencing. The low incidence of RUBISCO sub-units in the ICF's strongly suggests that the bands observed in figure 4 all originate from the apoplast and are not due to contamination by major intracellular constituents from cellular damage or leakage.

Analyses of the ICF's from tobacco leaves from different stages of development have revealed some interesting results. Polypeptide banding patterns of the ICF's from immature and mature leaf tissues were indicating that at particular different. verv developmental stages unique complements of proteins are present (figure 4). In situ staining for SOD enzyme activity has revealed a class of unique isoforms of SOD in the extracellular compartment. These bands represent a basic complement absent from the intracellular fraction. This result confirms the and extends the findings of Schinkel et al. (1998) who used vacuum infiltration to investigate the distribution of SOD in pine needles.

The large-scale variant of VI described here now provides the opportunity to isolate these unique complements of enzymes and proteins for further study.

We have successfully adapted this large-scale VI technique to barley and oil seed rape leaf materials and are currently exploring its potential application to We have successfully other plant tissue types. obtained several N-terminal sequences of apoplastic proteins from tobacco and other species, using this method (data not shown). Both time of exposure to vacuum and centrifugation speed to recover ICF's influence the quality and quantity of recovered from vacuum infiltrated tissue. It is clear that the best results are achieved only after some optimisation of the vacuum and centrifugation conditions for each experimental tissue. The methods described here should suffice for successful application to most plant materials. A further aspect of the methods versatility lies in the fact that many different types of buffer can be used to selectively release different fractions of apoplastic proteins (cf. Robertson et al 1998).

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Plant Molecular Biology 27, 59-67

Robertson, D, Davies, DR, Gerrish, C, Jupe, S, Bolwell, GP (1995)

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The plant peroxidase superfamily: Biochemical, molecular and physiological aspects of plant peroxidases University of Geneva pp3-13

# **APPENDIX 2**

**BLAST** search results

.

# **ITL4 - ATFDIVNQTYTVWA**

BLASTP 2.0.14 [Jun-29-2000]

Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402. RID: 963074119-21743-13743 Querv= (13 letters) Database: nr 517,398 sequences; 162,338,283 total letters >sp|P07052|PRR2\_TOBAC PATHOGENESIS-RELATED PROTEIN R MINOR FORM PRECURSOR (PR-R) (PROB12) (THAUMATIN-LIKE PROTEIN E2) pir/JJH0231 thaumatin-like protein E2 - common tobacco emb|CAA33292.1| (X15223) thaumatin-like protein [Nicotiana tabacum] emb|CAA27548.1| (X03913) TMV induced protein precursor (aa 1-226) [Nicotiana tabacum] prf||1206322A protein,TMV induced [Nicotiana sp ] Length = 226 Score = 27.0 bits (58), Expect = 25 Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ATFDIVNQDTYTV 13 ATFDIVNQ TYTV Sbjct: 26 ATFDIVNQCTYTV 38 \*\*\*\*\* >sp|P13046|PRR1\_TOBAC PATHOGENESIS-RELATED PROTEIN R MAJOR FORM PRECURSOR (THAUMATIN-LIKE PROTEIN E22) pir||JH0230 pathogenesis-related protein R precursor - common tobacco emb|CAA33293.1| (X15224) thaumatin-like protein [Nicotiana tabacum] emb|CAA31235.1| (X12739) pathogenesis-related protein R (AA 1 - 226) [Nicotiana tabacum] Length = 226 Score = 25.4 bits (54), Expect = 75 Identities = 11/13 (84%), Positives = 12/13 (91%) Query: 1 ATFDIVNQDTYTV 13 ATEDIVN+ TYTV Sbjct: 26 ATFDIVNKCTYTV 38 >emblCAB85637.1] (AJ237999) putative thaumatin-like protein [Vitis vinifera] Length = 222Score = 24.3 bits (51), Expect = 168 Identities = 10/13 (76%), Positives = 12/13 (91%) Query: 1 ATFDIVNQDTYTV 13 ATFDI+N+ TYTV Sbjct: 25 ATFDILNKCTYTV 37

\*\*\*\*\* >gb|AAB61590.1| (AF003007) VVTL1 [Vitis vinifera] Length = 222Score = 24.3 bits (51), Expect = 168 Identities = 10/13 (76%), Positives = 12/13 (91%) Query: 1 ATFDIVNQDTYTV 13 ATFDI+N+ TYTV Sbjct: 25 ATFDILNKCTYTV 37 >pir||PC2253 antifungal 27K protein - Diospyros texana (fragment) gb|AAB31495.1| antifungal protein thaumatin homolog {N-terminal} [Diospyros texana, overripe fruits, Peptide Partial, 30 aa] Length = 30 Score = 23.9 bits (50), Expect = 220 Identities = 10/13 (76%), Positives = 11/13 (83%) Query: 1 ATFDIVNQDTYTV 13 ATFDI N+ TYTV Sbjct: 1 ATFDIQNKXTYTV 13 >pir||T04212 osmotin precursor - Arabidopsis thaliana emb|CAB39936.1| (AL049500) osmotin precursor [Arabidopsis thaliana] emb[CAB78208.1] (AL161532) osmotin precursor [Arabidopsis thaliana] Length = 244 Score = 22.7 bits (47), Expect = 494 Identities = 9/13 (69%), Positives = 12/13 (92%) Query: 1 ATFDIVNQDTYTV 13 ATF+I+NQ +YTV

Sbjct: 23 ATFEILNQCSYTV 35

.

1: PC2253 . antifungal 27K pro...[gi:1083849]PubMed, Related Sequences Ο LOCUS 30 aa PLN 24-FEB-1995 PC2253 DEFINITION antifungal 27K protein - Diospyros texana (fragment). ACCESSION PC2253 g1083849 PID VERSION PC2253 GI:1083849 DBSOURCE pir: locus PC2253;

PubMedNucleotideProteinGenomeStructurePopSet Search for Preview/Index History Clipboard

summary: #length 30 #checksum 6033; PIR dates: 24-Feb-1995 #sequence\_revision 24-Feb-1995 #text\_change 24-Feb-1995:

punctuation in sequence.

#### KEYWORDS

Diospyros texana. SOURCE

**ORGANISM** Diospyros texana

Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; euphyllophytes; Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Asteridae; Ericales; Ebenaceae; Diospyros.

REFERENCE 1 (residues 1 to 30)

AUTHORS Vu,L. and Huynh,Q.K.

TITLE Isolation and characterization of a 27-kDa antifungal protein from the fruits of Diospyros texana

JOURNAL Biochem. Biophys. Res. Commun. 202 (2), 666-672 (1994) MEDLINE 94324951

COMMENT This protein belongs to one type of antifungal proteins called thaumatin-like protein due to the high degree of sequence homology to the thaumatin from Thaumatococcus danielli. Location/Qualifiers

FEATURES

source 1..30

/organism="Diospyros texana"

/db\_xref="taxon:37488" 1..30

Protein /product="antifungal 27K protein"

ORIGIN

1 atfdiqnkxt ytvwaaawap sypggxkqld

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II
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Revised: June 5, 2000.

Disclaimer | Write to the Help Desk NCBI | NLM | NIH

#### ITL 5 - SYQDFCVADD

BLASTP 2.0.14 [Jun-29-2000] Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402. RID: 963077133-10221-15128 Querv= (10 letters) Database: nr 517,398 sequences; 162,338,283 total letters Taxonomy reports Alignments >gb|AAB51571.1| (U75193) germin-like protein [Arabidopsis thaliana] gb|AAB51573.1| (U75195) germin-like protein [Arabidopsis thaliana] emb|CAA73213.1| (Y12673) GLP3 protein [Arabidopsis thaliana] dbj|BAA77208.1| (D89374) germin-like protein 2 precursor [Arabidopsis thaliana] emblCAB54516.1| (AJ132237) GER3 protein [Arabidopsis thaliana] Length = 211 Score = 32,4 bits (69), Expect = 0.22 Identities = 9/9 (100%), Positives = 9/9 (100%) Query: 1 SVQDFCVAD 9 SVQDFCVAD Sbjct: 21 SVQDFCVAD 29 >gb|AAB51241.1| (U81163) auxin-binding protein [Prunus persica] Length = 209 Score = 32.4 bits (69), Expect = 0.22 Identities = 9/9 (100%), Positives = 9/9 (100%) Query: 1 SVQDFCVAD 9 SVQDFCVAD Sbict: 19 SVQDFCVAD 27 >gb|AAD00295.1| (U79114) auxin-binding protein ABP19 [Prunus persica] Length = 209 Score = 32.4 bits (69), Expect = 0.22 Identities = 9/9 (100%), Positives = 9/9 (100%) Query: 1 SVQDFCVAD 9 SVQDFCVAD Sbjct: 19 SVQDFCVAD 27 >qb|AAB51566.1| (U75188) germin-like protein [Arabidopsis thaliana] gb|AAB51581.1| (U75203) germin-like protein [Arabidopsis thaliana] Length = 211 Score = 32.4 bits (69), Expect = 0.22 Identities = 9/9 (100%), Positives = 9/9 (100%) Query: 1 SVQDFCVAD 9 SVQDFCVAD Sbjct: 21 SVQDFCVAD 29  >sp|P45854|GLP1\_SINAL GERMIN-LIKE PROTEIN 1 PRECURSOR pir||T10454 germin-like protein 1 - white mustard emb|CAA59257.1| (X84786) Glp1 [Sinapis alba] Length = 211 Score = 32.4 bits (69), Expect = 0.22 Identities = 9/9 (100%), Positives = 9/9 (100%) Query: 1 SVQDFCVAD 9 SVQDFCVAD Sbjct: 21 SVQDFCVAD 29 >gb|AAB51750.1| (U95034) germin-like protein [Arabidopsis thaliana] Length = 204Score = 29.9 bits (63), Expect = 1.3 Identities = 8/9 (88%), Positives = 9/9 (99%) Query: 1 SVQDFCVAD 9 SVQDFCVA+ Sbjct: 14 SVQDFCVAN 22 \*\*\*\*\*\* >sp|P46271|GLP1\_BRANA GERMIN-LIKE PROTEIN PRECURSOR pir||T07854 germin-like protein (clone BnC4) - rape gb|AAA86365.1| (U21743) germin-like protein [Brassica napus] Length = 207 Score = 29.9 bits (63), Expect = 1.3 Identities = 8/9 (88%), Positives = 9/9 (99%) Query: 1 SVQDFCVAD 9 SVQDFCVA+ Sbjct: 18 SVQDFCVAN 26 >emb|CAA63014.1| (X91921) germin1 [Arabidopsis thaliana] Length = 208 Score = 29.9 bits (63), Expect = 1.3 Identities = 8/9 (88%), Positives = 9/9 (99%) Query: 1 SVQDFCVAD 9 SVQDFCVA+ Sbjct: 18 SVQDFCVAN 26 >gb|AAB51240.1| (U81162) auxin-binding protein [Prunus persica] Length = 214 Score = 29.9 bits (63), Expect = 1.3 Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 2 VQDFCVAD 9 VQDFCVAD Sbjct: 25 VQDFCVAD 32 

#### SICF2 - MPVDITYLQSAVRKGRVPL

BLASTP 2.0.10 [Aug-26-1999] Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402. Query= (19 letters) Database: Non-redundant GenBank CDS translations+PDB+SwissProt+SPupdate+PIR 416,691 sequences; 127,703,076 total letters emb|CAA18628.1| (AL022580) putative pectinacetylesterase protein [Arabidopsis thaliana] Length = 362 Score = 30.7 bits (65), Expect = 0.56 Identities = 10/11 (90%), Positives = 10/11 (90%) Query: 5 ITYLQSAVRKG 15 ITYLQSAV KG Sbjct: 26 ITYLQSAVAKG 36 spjQ62137|JAK3\_MOUSE TYROSINE-PROTEIN KINASE JAK3 (JANUS KINASE 3) (JAK-3) >gi|2137461|pir||158401 JAK3 - mouse >gi|537607 (L33768) JAK3 [Mus musculus] Length = 1299 Score = 28.2 bits (59), Expect = 3.3 Identities = 10/12 (83%), Positives = 10/12 (83%) Query: 8 LQSAVRKGRVPL 19 L SAV KGRVPL Sbjct: 452 LSSAVIKGRVPL 463 gb|AAD36370.1|AE001785\_1 (AE001785) ribonuclease H-related protein [Thermotoga maritima] Length = 223 Score = 27.3 bits (57), Expect = 5.9 Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 11 AVRKGRVP 18 AVRKGRVP Sbjct: 9 AVRKGRVP 16 

#### SICF3 - ASNGLGRTPQMGN

BLASTP 2.0.10 [Aug-26-1999] Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402. Query= (13 letters) Database: Non-redundant GenBank CDS translations+PDB+SwissProt+SPupdate+PIR emb|CAB00120.1| (Z75955) similar to alpha-N-acetylgalactosaminidase; cDNA EST EMBL:M92935 comes from this gene; cDNA EST CEMSA04R comes from this gene [Caenorhabditis elegans] Length = 451 Score = 30.3 bits (64), Expect = 0.76 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 3 NGLGRTPQMG 12 NGLGRTP MG Sbjct: 18 NGLGRTPPMG 27 emb|CAA64760| (X95506) alpha-galactosidase MEL [Saccharomyces sp.] Length = 471 Score = 28.2 bits (59), Expect = 3.3 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 3 NGLGRTPQMG 12 NGLG TPQMG Sbjct: 24 NGLGLTPQMG 33 pir||JQ1021 alpha-galactosidase (EC 3.2.1.22) - yeast (Saccharomyces cerevisiae) >gi|171924 (M58484) alpha-galactosidase [Saccharomyces carlsbergensis] Length = 471 Score = 28.2 bits (59), Expect = 3.3 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 3 NGLGRTPQMG 12 NGLG TPQMG Sbjct: 24 NGLGLTPQMG 33 sp|P41947|MEL6\_YEAST ALPHA-GALACTOSIDASE 6 PRECURSOR (MELIBIASE 6) (ALPHA-D-GALACTOSIDE GALACTOHYDROLASE 6) >gi|1084809|pir||S50312 MEL6 protein - yeast (Saccharomyces cerevisiae) >gi|547466|emb|CAA85739| (Z37510) alpha-galactosidase [Saccharomyces cerevisiae] Length = 471 Score = 28.2 bits (59), Expect = 3.3 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 3 NGLGRTPQMG 12 NGLG TPQMG Sbjct: 24 NGLGLTPQMG 33  Other Formats: <Picture: [FASTA Format]> <Picture: [Graphical View]> Links: <Picture: [1 medline link]> <Picture: [1 nucleotide link]> <Picture: [92 protein neighbors]>

INV 02-SEP-1999 CAB00120 451 aa LOCUS DEFINITION similar to alpha-N-acetylgalactosaminidase; cDNA EST EMBL:M92935 comes from this gene; cDNA EST CEMSA04R comes from this gene. ACCESSION CAB00120 g3878996 PID VERSION CAB00120.1 GI:3878996 DBSOURCE embl locus CER07B7, accession Z75955.1 **KEYWORDS** SOURCE Caenorhabditis elegans. ORGANISM Caenorhabditis elegans Eukaryota; Metazoa; Nematoda; Secernentea; Rhabditia; Rhabditida; Rhabditina; Rhabditoidea; Rhabditidae; Peloderinae; Caenorhabditis. REFERENCE 1 (residues 1 to 451) AUTHORS Wilson, R., Ainscough, R., Anderson, K., Baynes, C., Berks, M., Bonfield, J., Burton, J., Connell, M., Copsey, T., Cooper, J., Coulson,A., Craxton,M., Dear,S., Du,Z., Durbin,R., Favello,A., Fulton,L., Gardner,A., Green,P., Hawkins,T., Hillier,L., Jier,M., Johnston, L., Jones, M., Kershaw, J., Kirsten, J., Laister, N., Latreille, P., Lightning, J., Lloyd, C., McMurray, A., Mortimore, B., O'Callaghan, M., Parsons, J., Percy, C., Rifken, L., Roopra, A., Saunders, D., Shownkeen, R., Smaldon, N., Smith, A., Sonnhammer, E., Staden, R., Sulston, J., Thierry-Mieg, J., Thomas, K., Vaudin, M., Vaughan, K., Waterston, R., Watson, A., Weinstock, L., Wilkinson-Sproat, J. and Wohldman, P. TITLE 2.2 Mb of contiguous nucleotide sequence from chromosome III of C. elegans JOURNAL Nature 368 (6466), 32-38 (1994) MEDLINE 94150718 REFERENCE 2 (residues 1 to 451) AUTHORS Harris, B. TITLE Direct Submission JOURNAL Submitted (13-JUL-1996) Louis, MO 63110, USA. E-mail: jes@sanger.ac.uk or rw@nematode.wustl.edu Coding sequences below are predicted from computer analysis, using COMMENT predictions from Genefinder (P. Green, U. Washington), and other available information. For a graphical representation of this sequence and its analysis see'http://webace.sanger.ac.uk/cgibin/display?db=wormace&class=Sequence &object=R07B7 Current sequence finishing criteria for the C. elegans genome sequencing consortium are that all bases are either sequenced unambiguously on both strands, or on a single strand with both a dye primer and dye terminator reaction, from distinct subclones. Exceptions are indicated by an explicit note. IMPORTANT: This sequence is NOT necessarily the entire insert of the specified clone. It may be shorter because we only sequence overlapping sections once, or longer because we arrange for a small overlap between neighbouring submissions. This sequence is the entire insert of clone R07B7. The true right end of clone F10C2 is at 1275 in this sequence. The start of this sequence (1..112) overlaps with the end of sequence Z81497. The end of this sequence (43364..43469) overlaps with the start of sequence Z92825. FEATURES Location/Qualifiers 1..451 source /organism="Caenorhabditis elegans" /db xref="taxon:6239" /chromosome="V" /clone="R07B7" 1..451 Protein /name="similar to alpha-N-acetylgalactosaminidase; cDNA EST EMBL:M92935 comes from this gene; cDNA EST CEMSA04R comes from this gene" 1..451 CDS /gene="R07B7.11" /db xref="SPTREMBL:Q21801" /coded by="join(Z75955.1:28833..29013, Z75955.1:29059..29198,Z75955.1:29270..29899, Z75955.1:29950..30051,Z75955.1:30120..30422)"

#### ORIGIN

RIGIN 1 mrlllpllfc vgafcldngl grtppmgwms wtafyceidc vkhptgcine qlykdmadql 61 vsggydkvgy ksvhiddcws emerdshgil vanktrfpsg mkalakymhd rglkfgiyed 121 ygtktcggyp gsykhekvda qtfaawdvdy lkldgcnidq ammpigyplf ekelnetgrp 181 imyscswpay lidhpelvny nligkycntw rnfddinssw ksiisiisyy dkmqdkhipt 241 hgpgkwhdpd mlvignkgit Idmsisqftv wciwsaplim sndlriigds fkdvlknkea 301 ikinqdplgi mgrliknstd igvyvkqitp skgdkksfaf aylnrnekeg ykrieiqlas 361 igltdpagyy vhdiwshvdl gllrsgdsiv vsiapagsvf fradlaslhq pkgkrrykfd 421 nsnvvyefer eqsqlgrfdf gdylrtnrvq f

No significant results

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# SICF8 - AVLDFCVGDLSVPDNPAG

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TBLASTN 2.0.10 [Aug-26-1999] Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402. Querv= (18 letters) Database: Non-redundant GenBank+EMBL+DDBJ+PDB sequences 478,564 sequences; 1,310,507,524 total letters If you have any problems or questions with the results of this search please refer to the BLAST FAQs gb|U81162|PPU81162 Prunus persica auxin-binding protein (ABP20) mRNA, complete cds Length = 1001Score = 37.1 bits (80), Expect = 0.026 Identities = 12/18 (66%), Positives = 14/18 (77%) Frame = +1 Query: 1 AVLDFCVGDLSVPDNPAG 18 AV DFCV DL+ P+ PAG Sbjct: 103 AVQDFCVADLAAPEGPAG 156 emb|Y15962|HVGLP1 Hordeum vulgare mRNA for germin-like protein 1 Length = 833Score = 33.3 bits (71), Expect = 0.37 Identities = 11/15 (73%), Positives = 11/15 (73%) Frame = +1 Query: 4 DFCVGDLSVPDNPAG 18 DFCV DLS D PAG Sbict: 106 DFCVADLSCSDTPAG 150 dbj|AB010876|AB010876 Oryza sativa OsGLP1 mRNA for germin-like protein 1, complete cds Length = 994 Score = 30.7 bits (65), Expect = 2.2 Identities = 10/15 (66%), Positives = 11/15 (72%) Frame = +1 Query: 4 DFCVGDLSVPDNPAG 18 DFCV DL+ D PAG Sbjct: 181 DFCVADLTCSDTPAG 225 gb|AF032975|AF032975 Oryza sativa germin-like protein 5 (GER5) mRNA, complete cds Length = 1030 Score = 30.7 bits (65), Expect = 2.2 Identities = 10/15 (66%), Positives = 11/15 (72%) Frame = +2Query: 4 DFCVGDLSVPDNPAG 18 DFCV DL+ D PAG Sbict: 182 DFCVADLTCSDTPAG 226

Other Formats: <Picture: [FASTA Format]> <Picture: [Graphical View]> Links: <Picture: [1 nucleotide link]> <Picture: [105 protein neighbors]> LOCUS AAB51240 214 aa PLN 02-APR-1997 DEFINITION auxin-binding protein. ACCESSION AAB51240 PID g1916807 VERSION AAB51240.1 GI:1916807 DBSOURCE locus PPU81162 accession U81162.1 KEYWORDS peach. SOURCE **ORGANISM** Prunus persica Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; euphyllophytes; Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae; eurosids I; Rosales; Rosaceae; Prunus. REFERENCE 1 (residues 1 to 214) AUTHORS Ohmiya, A., Tanaka, Y., Kadowaki, K. and Hayashi, T. TITLE Cloning of genes encoding auxin-binding proteins (ABP19/20) from peach JOURNAL Unpublished REFERENCE 2 (residues 1 to 214) AUTHORS Ohmiya,A., Tanaka,Y., Kadowaki,K. and Hayashi,T. TITLE Direct Submission JOURNAL Submitted (05-DEC-1996) Department of Breeding, National Institute of Fruit Tree Science, Fujimoto 2-1, Tsukuba, Ibaraki 305, Japan COMMENT Method: conceptual translation. Location/Qualifiers FEATURES 1..214 source /organism="Prunus persica" /cultivar="Akatsuki" /db\_xref="taxon:3760" 1..214 Protein /product="auxin-binding protein" /name="cell wall protein" CDS 1..214 /aene="ABP20" /coded\_by="U81162.1:34..678" ORIGIN 1 mpqatmifpi lftfflllss snaavqdfcv adlaapegpa gfsckkpasv kvndfvfsgl

61 giagntsnii kaavtpafva qfpgvnglgi siarldlavg gvvpfhthpg asevlivaqg

121 ticagfvasd ntpylqtlek gdimvfpqgl lhfqvnggea palafasfgs aspglqildf

181 alfkndlpte viaqttflda aqikklkgvl ggtn

<Picture: Entrez protein Query>

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# SICF9 - AVLDFC<

Sequencing stopped because of similarity to SICF8.

See sequence search results for SICF8



# SICF11 – MPDP?PTFAE

No homologies found